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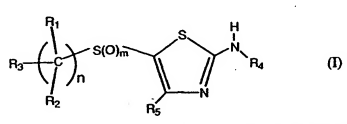
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(54) Title: AMINOTHIAZOLE INHIBITORS OF CYCLIN DEPENDENT KINASES



(57) Abstract: Compounds of formula (I) and pharmaceutically acceptable salts thereof. As used in formula (I), and throughout the specification, the symbols have the following meanings: R₁ and R₂ are independently hydrogen, fluorine or alkyl; R₃ is aryl or heteroaryl, R₄ has various meanings; R₅ is hydrogen or alkyl; m is an integer of 0 to 2; and n is an integer of 1 to 3. The compounds of formula (I) are pro-

tein kinase inhibitors and are useful in the treatment and prevention of proliferative diseases, for example, cancer, inflammation and arthritis. They may also be useful in the treatment of neurodegenerative diseases such as Alzheimer's diseases cardiovascular diseases, viral diseases and fungal diseases.

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AMINOTHIAZOLE INHIBITORS OF CYCLIN DEPENDENT KINASES

Brief Description of the Invention

The present invention is directed to compounds of the formula

$$R_3 \xrightarrow{R_1} S(O)_m \xrightarrow{S} N \xrightarrow{H} R_4 \qquad (I)$$

and pharmaceutically acceptable salts thereof. As used in formula I, and throughout the specification, the symbols have the following meanings:

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

R₃ is aryl or heteroaryl

R4 is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl,

heteroaryl, heteroarylalkyl, heterocycloalkyl,

heterocycloalkylalkyl; or

CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl,

 $CO\text{-}heteroaryl, CO\text{-}alkyl\text{-}heteroaryl, CO\text{-}heterocycloalkyl,}$

CO-alkyl-heterocycloalkyl; or

CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl,

CONH-alkyl-aryl, CONH-heteroaryl,

CONH-alkyl-heteroaryl, CONH-heterocycloalkyl,

20 CONH-alkyl-heterocycloalkyl; or

COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl,

COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl,

COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or

 SO_2 -cycloalkyl, SO_2 -aryl, SO_2 -alkyl-cycloalkyl, SO_2 -alkyl-aryl,

SO₂-heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl,

 SO_2 -alkyl-heterocycloalkyl; or

C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl, C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl,

C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl,
C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or
C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl,
C(NNO₂)NH-alkyl-cycloalkyl, C(NNO₂)NH-alkyl-aryl,
C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl,
C(NNO₂)NH-heterocyloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl;

or

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C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl, C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl, C(NH)NH-alkyl-heteroaryl, C(NH)NH-alkyl-heteroaryl,

C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,

C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl, C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl,

15 C(NH)NHCO-heterocylcloalkyl,

C(NH)NHCO-alkyl-heterocycloalkyl; or

C(NOR6)NH-alkyl, C(NOR6)NH-cycloalkyl, C(NOR6)NH-aryl,

C(NOR₆)NH-alkyl-cycloalkyl, C(NOR₆)NH-alkyl-aryl,

C(NOR₆)NH-heteroaryl, C(NOR₆)NH-alkyl-heteroaryl,

20 C(NOR₆)NH-heterocylcoalkyl, C(NOR₆)NH-alkyl-heterocycloalkyl;

R₅ is hydrogen or alkyl;

R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

m is an integer of 0 to 2; and n is an integer of 1 to 3.

The compounds of formula I are protein kinase inhibitors and are useful in the treatment and prevention of proliferative diseases, for example, cancer, inflammation and arthritis. They may also be useful in the treatment of neurodegenerative diseases such as Alzheimer's disease, cardiovascular diseases, viral diseases and fungal diseases.

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Description of the Invention

The present invention provides for compounds of formula I,
pharmaceutical compositions employing such compounds and for methods
of using such compounds.

Listed below are definitions of various terms used to describe the compounds of the instant invention. These definitions apply to the terms as they are used throughout the specification (unless they are otherwise limited in specific instances) either individually or as part of a larger group.

It should be noted that any heteroatom with unsatisfied valances is assumed to have the hydrogen atom to satisfy the valances.

Carboxylate anion refers to a negatively charged group -COO .

The term "alkyl" or "alk" refers to a monovalent alkane (hydrocarbon) derived radical containing from 1 to 12 carbon atoms unless otherwise defined. An alkyl group is an optionally substituted straight, branched or cyclic saturated hydrocarbon group. When substituted, alkyl groups may be substituted with up to four substituent groups, R as defined, at any available point of attachment. When the alkyl group is said to be substituted with an alkyl group, this is used interchangeably with "branched alkyl group". Exemplary unsubstituted such groups include methyl, ethyl, propyl, isopropyl, n-butyl, t-butyl, isobutyl, pentyl, hexyl, isohexyl, heptyl, 4,4-dimethylpentyl, octyl, 2,2,4-trimethylpentyl, nonyl, decyl, undecyl, dodecyl, and the like. Exemplary substituents may include but are not limited to one or more of the following groups: halo (such as F, Cl, Br, I), haloalkyl (such as CCl3 or CF3), alkoxy, alkylthio, hydroxy, carboxy (-COOH), alkyloxycarbonyl (-C(O)R), alkylcarbonyloxy (-OCOR), amino (-NH2), carbamoyl

30 (-NHCOOR- or -OCONHR-), urea (-NHCONHR-) or thiol (-SH). Alkyl

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groups as defined may also comprise one or more carbon to carbon double bonds or one or more carbon to carbon triple bonds.

The term "alkenyl" refers to a hydrocarbon radical straight, branched or cyclic containing from 2 to 12 carbon atoms and at least one carbon to carbon double bond.

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The term "alkynyl" refers to a hydrocarbon radical straight, branched or cyclic containing from 2 to 12 carbon atoms and at least one carbon to carbon triple bond.

Cycloalkyl is a specie of alkyl containing from 3 to 15 carbon atoms, without alternating or resonating double bonds between carbon atoms. It may contain from 1 to 4 rings. Exemplary unsubstituted such groups include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, adamantyl, etc. Exemplary substituents include one or more of the following groups: halogen, alkyl, alkoxy, alkyl hydroxy, amino, nitro, cyano, thiol and/or alkylthio.

The terms "alkoxy" or "alkylthio", as used herein, denote an alkyl group as described above bonded through an oxygen linkage (-O-) or a sulfur linkage (-S-), respectively.

The term "alkyloxycarbonyl", as used herein, denotes an alkoxy group bonded through a carbonyl group. An alkoxycarbonyl radical is represented by the formula: -C(O)OR, where the R group is a straight or branched C₁₋₆ alkyl group.

The term "alkylcarbonyl" refers to an alkyl group bonded through a carbonyl group.

The term "alkylcarbonyloxy", as used herein, denotes an alkylcarbonyl group which is bonded through an oxygen linkage.

The term "arylalkyl", as used herein, denotes an aromatic ring bonded to an alkyl group as described above.

The term "aryl" refers to monocyclic or bicyclic aromatic rings, e.g. phenyl, substituted phenyl and the like, as well as groups which are fused, e.g., napthyl, phenanthrenyl and the like. An aryl group thus contains at

least one ring having at least 6 atoms, with up to five such rings being present, containing up to 22 atoms therein, with alternating (resonating) double bonds between adjacent carbon atoms or suitable heteroatoms. Aryl groups may optionally be substituted with one or more groups including, but not limited to halogen, alkyl, alkoxy, hydroxy, carboxy, carbamoyl, alkyloxycarbonyl, nitro, trifluoromethyl, amino, cycloalkyl, cyano, alkyl S(O)_m (m=O, 1, 2), or thiol.

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The term "heteroaryl" refers to a monocyclic aromatic hydrocarbon group having 5 or 6 ring atoms, or a bicyclic aromatic group having 8 to 10 atoms, containing at least one heteroatom, O, S, or N, in which a carbon or nitrogen atom is the point of attachment, and in which one or two additional carbon atoms is optionally replaced by a heteroatom selected from O or S, and in which from 1 to 3 additional carbon atoms are optionally replaced by nitrogen heteroatoms, said heteroaryl group being optionally substituted as described herein. Exemplary heteroaryl groups include the following: thienyl, furyl, pyrrolyl, pyridinyl, imidazolyl, pyrrolidinyl, piperidinyl, thiazolyl, oxazolyl, triazolyl, pyrazolyl, isoxazolyl, isothiazolyl, pyrazinyl, pyridazinyl, pyrimidinal, triazinylazepinyl, indolyl, isoindolyl, quinolinyl, isoquinolinyl, benzothiazolyl, benzoxazolyl, benzimidazolyl, benzoxadiazolyl, benzofurazanyl and tetrahydropyranyl. Exemplary substituents include one or more of the following: halogen, alkyl, alkoxy, hydroxy, carboxy, carbamoyl, alkyloxycarbonyl, trifluoromethyl, cycloalkyl, nitro, cyano, amino, alkylS(O)_m (m=0, 1, 2), or thiol.

The term "heteroarylium" refers to heteroaryl groups bearing a quaternary nitrogen atom and thus a positive charge.

The term "heterocycloalkyl" refers to a cycloalkyl group (nonaromatic) in which one of the carbon atoms in the ring is replaced by a heteroatom selected from O, S or N, and in which up to three additional carbon atoms may be replaced by said heteroatoms.

The term "quaternary nitrogen" refers to a tetravalent positively charged nitrogen atom including, e.g. the positively charged nitrogen in a tetraalkylammonium group (e.g. tetramethylammonium, N-methylpyridinium), the positively charged nitrogen in protonated ammonium species (e.g. trimethylhydroammonium, N-hydropyridinium), the positively charged nitrogen in amine N-oxides (e.g. N-methyl-morpholine-N-oxide, pyridine -N-oxide), and the positively charged nitrogen in an N-amino-ammonium group (e.g.

N-aminopyridinium).

The term "heteroatom" means O, S or N, selected on an

independent basis.

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The term "halogen" or "halo" refers to chlorine, bromine, fluorine or iodine.

When a functional group is termed "protected", this means that the group is in modified form to preclude undesired side reactions at the protected site. Suitable protecting groups for the compounds of the present invention will be recognized from the present application taking into account the level of skill in the art, and with reference to standard textbooks, such as Greene, T. W. et al., *Protective Groups in Organic Synthesis*, Wiley, N.Y. (1991).

Suitable examples of salts of the compounds according to the invention with inorganic or organic acids are hydrochloride, hydrobromide, sulfate, phosphate. Salts which are unsuitable for pharmaceutical uses but which can be employed, for example, for the isolation or purification of free compounds I or their pharmaceutically acceptable salts, are also included.

All stereoisomers of the compounds of the instant invention are contemplated, either in admixture or in pure or substantially pure form. The definition of the compounds according to the invention embraces all possible stereoisomers and their mixtures. It very particularly embraces the racemic forms and the isolated optical isomers having the specified

activity. The racemic forms can be resolved by physical methods, such as, for example, fractional crystallization, separation or crystallization of diastereomeric derivatives or separation by chiral column chromatography. The individual optical isomers can be obtained from the racemates by conventional methods, such as, for example, salt formation with an optically active acid followed by crystallization.

It should be understood that solvates (e.g., hydrates) of the compounds of formula I are also within the scope of the present invention. Methods of solvation are generally known in the art. Accordingly, the compounds of the instant invention may be in the free or hydrate form, and may be obtained by methods exemplified by the following schemes.

Scheme 1

DTT HS
$$NH_2$$
 $R_3(CR_1R_2)_n$ -L $R_3(R_2R_1C)_n$ NHR_4

(V) (I)

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As illustrated in Scheme 1, compounds of formula I where X is S are prepared by reacting 2-aminothiazole (II) with bromine in the presence of sodium or potassium thiocyanate to obtain a thiocyanated aminothiazole, specifically 5-thiocyanatoaminothiazole (III). Compound III is then reacted with R₄-L, where L is a leaving group such as a halogen, in the presence of a base such as triethylamine to provide a 5-thiocyanatothiazole intermediate (IV), where R₄ is as defined in the specification. The intermediate (IV) is then reduced to a thiol (V) using

reducing agents such as dithiothreitol (DTT), sodium borohydride, zinc or other known reducing agents. Compound (V) is then reacted with alkyl, aryl or heteroaryl halides, such as R_3 (CR_1R_2)_n-L, where L is a leaving group such as a halogen, in the presence of a base such as potassium carbonate to obtain compounds of formula I. The steps of reducing the thiocyanothiazole intermediate (IV) to the thiol (V), and the reaction of the reduced thiol (V) to provide compounds of formula I where X is S, may be carried out sequentially without purification.

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Scheme 2

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In Scheme 2, 5-thioacetyl-2-acetylaminothiazole of structure VI is reacted with an alkoxide such as potassium t-butoxide in alcohol or THF solvent and the resulting thiol is reacted in situ with a group of formula $R_3(CR_1R_2)_n$ -L (where L is a leaving group, such as a halogen) such as 2-halomethyloxazole (VII) to provide a compound such as formula VIII, wherein R_1 and R_2 are hydrogen, and R_6 is acetyl. The 2-halomethyloxazole compounds of formula VII may be prepared using

several synthetic routes known in the art. Chem. Pharm. Bull. 30, 1865 (1982); Bull. Chem. Soc. Japan (52, 3597 (1979); JCS Chem. Comm. 322 (1981); Comprehensive Heterocyclic Chemistry, vol. 6, 177, edited by A. Katritzky and C.W. Rees, Pergamon Press (1984).

Compounds of formula VIII (a compound of formula I where R₄ is acetyl and X is sulfur) can be hydrolyzed in the presence of a base such as sodium hydroxide to provide a compound of formula IX. A compound of formula IX may then be reacted with R₄-L, in the presence of a base such as triethylamine, where L is a leaving group such as a halogen, to give compounds of formula I where X is sulfur. In this manner, compounds of formula IX, which is a compound of formula I where R₄ is hydrogen, can be treated with agents such as isothiocyanates, halides, acyl halides, chloroformates, isocyanates or sulfonyl chlorides to provide thioureas, amines, amides, carbamates, ureas or sulfonamides. The procedures in Scheme 2 specifically illustrate a methyloxazole group, but are general for all R₃(CR₁R₂)_n- groups specified by formula I.

Alternatively, compounds of formula VII, where L is bromine, may be prepared by halogenation of 2-methyloxazole using N-bromosuccinimide in the presence of dibenzoylperoxide.

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Scheme 3

Scheme 3

$$R_8 \stackrel{OH}{\longrightarrow} NH_2 + CI \stackrel{R_1}{\longrightarrow} R_2 \stackrel{Et_3N}{\longrightarrow} N$$
 (X)
 (X)
 (X)
 $R_8 \stackrel{H}{\longrightarrow} NH_2 \stackrel{R_1}{\longrightarrow} R_2 \stackrel{R_2}{\longrightarrow} CI$
 (X)
 (X)

Scheme 3 illustrates an alternative method of preparing compound VII, which is a compound of formula R₃(CR₁R₂)_n-L where L is chlorine and n is the integer l. In this scheme, compound VII is prepared by the reaction of a compound of formula X and formula XI in the presence of a base such as triethylamine to provide compounds of formula XII.

Compound XII may be oxidized by an oxidant such as oxalylchloride/DMSO in the presence of a base such as triethylamine to provide a compound of formula XIII which may be cyclized by an agent such as phosphorous oxychloride to provide compounds of formula VII, wherein L is chlorine. Alternatively, compounds of formula XIII may be prepared by reaction of the amino ketone correponding to X with an acid chloride such as XI.

Scheme 4 $R_{8} \stackrel{O}{\underset{R_{7}}{\bigvee}} R_{2} + NC \stackrel{R_{1}}{\underset{Cl}{\bigvee}} R_{2} \xrightarrow{BF_{3}.Et_{2}O} \stackrel{R_{7}}{\underset{R_{8}}{\bigvee}} R_{7} \stackrel{N}{\underset{R_{1}}{\bigvee}} Cl$ $(XIV) \qquad (XV) \qquad (XV)$

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Compounds of formula VII, where L is chlorine, may also be prepared from the reaction of diazoketones as illustrated by formula XIV in Scheme 4 with chloronitriles, such as indicated by formula XV, in the presence of BF₃ etherate to provide compounds of formula VII, wherein L is chlorine.

Scheme 5

(XXIII)

$$R_4$$
 TFA
 CH_2Cl_2
 $Step 7$
 R_1
 R_2
 R_1
 R_2
 R_1
 R_2
 R_3
 R_1
 R_2
 R_3

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In Scheme 5, starting compound XVI denotes a resin-bound benzyl alcohol support used for solid phase synthesis which is prepared from a Merrifield resin denoted as Q, and 2-methoxy-4-hydroxybenzaldehyde, followed by reduction with reducing agents such as NaBH₄. In step 1, starting compound XVI is treated with triphosgene and triphenylphosphine (PPh₃) in dichloromethane to give the chlorobenzyl resin of formula XVII. In step 2, a thiocyanato trifluoroacetamide (XVIII) is alkylated with the resin-bound benzyl chloride (XVII) in the presence of diisopropylethylamine (DIPEA) to form a resin-bound thiocyanate (XIX). The thiocyanato trifluoroacetamide compound of formula XVII is prepared by reacting 5-thiocyanatoaminothiazole of formula III (Scheme I) with trifluoroacetic anhydride using a base such as 2,6-lutidine.

The resin-bound thiocyanate (XIX) is then reduced to a resin-bound thiol (XX) in step 3 with reducing agent such as dithiothreitol (DTT) in tetrahydrofuran (THF) and methanol. The resulting resin-bound thiol 15 (XX) is reacted with R₃(CR₁R₂)_n-L, where L is a leaving group, in the presence of a base such as 1,8-diazabicyclo[5,4,0]undec-7-ene (DBU) at 80 °C in dimethylformamide (DMF) to form compounds of formula XXI (step 4). Deprotection of the trifluoroacetyl group of compound XXI is performed in step 5 using sodium borohydride to provide a compound of 20 formula XXII. In step 6, the deprotected compound XXII is reacted with R₆X, where X is a leaving group, in the presence of a base such as diisopropylethylamine to provide compounds of formula XXIII. The product is then cleaved from the solid phase resin in step 7 with trifluoroacetic acid (TFA) to give compounds of formula I where X is 25 sulfur. Compounds of formula I where X is $S(O)_m$ and m is 1 or 2 may be prepared from compounds of formula I where m is 0 by oxidation with an oxidant such as sodium periodate, meta-chloroperbenzoic acid, or oxone.

Scheme 6

5 Scheme 6 illustrates the preparation of compounds of formula I from a 2bromo thiazole XXIV. A compound of formula IX is reacted with a diazotizing agent such as tBuONO in the presence of copper bromide to provide the exemplary 2-bromo thiazole of formula XXIV. Compound XXIV may then be reacted with a compound of formula R₄NH₂, with or without an added base, to provide compounds of formula I.

Scheme 7

S Br
$$R_4NH_2$$
 S NHR_4 R_1R_2 NHR_4 R_1 R_2 NHR_4 R_1 R_2 R_1 R_2 R_2 R_3 R_4 R_4 R_5 R_6 R_8 R_1 R_2 R_8 R_8 R_1 R_2 R_8 R_1 R_2 R_2 R_2 R_1 R_2 R_2 R_2 R_2 R_2 R_3 R_1 R_2 R_2 R_3 R_1 R_2 R_2 R_2 R_3 R_1 R_2 R_3 R_2 R_3 R_1 R_2 R_3 R_2 R_3 R_1 R_2 R_3 R_2 R_3 R_3

Compounds of formula I may also be prepared starting from 2-bromothiazole XXV by reaction with a compound of formula R₄NH₂, with or without an added base, to provide a compound of formula XXVI. The compound of formula XXVI may be reacted with a thiocyanating agent such as sodium thiocyanate in the presence of bromine to provide a compound of formula IV, that may then be converted to a compound of formula I as described in Scheme 1. Alternatively, the compound of formula XXVI may be treated with a brominating agent such as bromine in acetic acid to generate a compound XXVII. Compounds of formula XXVII may be reacted with either XXVIII or XXIX (themselves available from a compound of formula VII) in the presence of base to provide compounds of formula I.

The starting compounds of Schemes 1-7 are commercially available or may be prepared by methods known to one of ordinary skill in the art.

All compounds of formula I may be prepared by modification of the procedures described herein.

The preferred compounds of formula I are those where:

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_7

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wherein Y is oxygen, sulfur or NR9;

R₄ is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl,

heterocycloalkylalkyl; or

25 CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl, CO-heteroaryl, CO-alkyl-heteroaryl, CO-heterocycloalkyl, CO-alkyl-heterocycloalkyl; or

CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl, CONH-alkyl-aryl, CONH-heteroaryl,

CONH-alkyl-heteroaryl, CONH-heterocycloalkyl, CONH-alkyl-heterocycloalkyl; or COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl, COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl, COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or 5 SO2-cycloalkyl, SO2-aryl, SO2-alkyl-cycloalkyl, SO2-alkyl-aryl, SO₂-heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl, SO₂-alkyl-heterocycloalkyl; or C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl, C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl, 10 C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl, C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl, C(NNO₂)NH-alkyl-cycloalkyl, C(NNO₂)NH-alkyl-aryl, C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl, 15 C(NNO₂)NH-heterocyloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl; or C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl, C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl, 20 C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl, C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl, C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl, C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl, C(NH)NHCO-heterocylcloalkyl, 25 C(NH)NHCO-alkyl-heterocycloalkyl; or C(NOR₆)NH-alkyl, C(NOR₆)NH-cycloalkyl, C(NOR₆)NH-aryl, C(NOR₆)NH-alkyl-cycloalkyl, C(NOR₆)NH-alkyl-aryl, C(NOR₆)NH-heteroaryl, C(NOR₆)NH-alkyl-heteroaryl, C(NOR₆)NH-heterocylcoalkyl, C(NOR₆)NH-alkyl-heterocycloalkyl; 30 R₅ is hydrogen; and

R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

R₇ and R₈ are independently hydrogen, alkyl, cycloalkyl, aryl, alkylcycloalkyl, alkylaryl, heteroaryl, alkylheteroaryl, heterocycloalkyl, alkylheterocycloalkyl or halogen;

Re is H or alkyl; m is the integer 0; and

n is the integer 1.

The most preferred compounds of formula I are those where:

R₁ is hydrogen;

R₂ is hydrogen, fluorine or alkyl;

R₃ is a substituted oxazole having the configuration:

$$- \bigvee_{N}^{\mathsf{R}_{\mathsf{B}}} \mathsf{R}_{\mathsf{7}}$$

15 R4 is CO-alkyl, CO-alkyl-aryl, CO-cycloalkyl,

CO-alkyl-heteroaryl, CO-alkyl-heteroalkyl,

CO-alkyl-heterocycloalkyl, aryl, arylalkyl, heteroaryl,

heteroarylalkyl;

R₅ is hydrogen;

20 R₇ is hydrogen;

R₈ is an alkyl group, such as tert-butyl;

m is the integer 0; and

n is the integer 1.

25 The compounds according to the invention have pharmacological properties; in particular, the compounds of formula I are inhibitors of protein kinases such as the cyclin dependent kinases (cdks), for example, cdc2 (cdk1), cdk2, and cdk4. The novel compounds of formula I are expected to be useful in the therapy of proliferative diseases such as

cancer, autoimmune diseases, viral diseases, fungal diseases, neurodegenerative disorders and cardiovascular disease.

More specifically, the compounds of formula I are useful in the treatment of a variety of cancers, including (but not limited to) the following:

-carcinoma, including that of the bladder, breast, colon, kidney, liver, lung, including small cell lung cancer, esophagus, gall bladder, ovary, pancreas, stomach, cervix, thyroid, prostate, and skin, including squamous cell carcinoma;

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-hematopoietic tumors of lymphoid lineage, including leukemia, acute lymphocytic leukemia, acute lymphoblastic leukemia, B-cell lymphoma, T-cell lymphoma, Hodgkins lymphoma, non-Hodgkins lymphoma, hairy cell lymphoma and Burkett's lymphoma;

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-hematopoietic tumors of myeloid lineage, including acute and chronic myelogenous leukemias, myelodysplastic syndrome and promyelocytic leukemia;

-tumors of mesenchymal origin, including fibrosarcoma and rhabdomyosarcoma;

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tumors of the central and peripheral nervous system,
 including astrocytoma, neuroblastoma, glioma and schwannomas; and

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-other tumors, including melanoma, seminoma, teratocarcinoma, osteosarcoma, xenoderoma pigmentosum, keratoctanthoma, thyroid follicular cancer and Kaposi's sarcoma.

Due to the key role of cdks in the regulation of cellular proliferation in general, inhibitors could act as reversible cytostatic agents which may be useful in the treatment of any disease process which features abnormal cellular proliferation, e.g., benign prostate hyperplasia, familial adenomatosis polyposis, neuro-fibromatosis, atherosclerosis, pulmonary

fibrosis, arthritis, psoriasis, glomerulonephritis, restenosis following angioplasty or vascular surgery, hypertrophic scar formation, inflammatory bowel disease, transplantation rejection, endotoxic shock, and fungal infections.

Compounds of formula I may also be useful in the treatment of Alzheimer's disease, as suggested by the recent finding that cdk5 is involved in the phosphorylation of tau protein (*J. Biochem*, 117, 741-749 (1995)).

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Compounds of formula I may induce or inhibit apoptosis. The apoptotic response is aberrant in a variety of human diseases. Compounds of formula I, as modulators of apoptosis, will be useful in the treatment of cancer (including but not limited to those types mentioned hereinabove), viral infections (including but not limited to herpevirus, poxvirus, Epstein-Barr virus, Sindbis virus and adenovirus), prevention of AIDS development in HIV-infected individuals, autoimmune diseases (including but not limited to systemic lupus, erythematosus, autoimmune mediated glomerulonephritis, rheumatoid arthritis, psoriasis, inflammatory bowel disease, and autoimmune diabetes mellitus), neurodegenerative disorders (including but not limited to Alzheimer's disease, AIDS-related dementia, Parkinson's disease, amyotrophic lateral sclerosis, retinitis pigmentosa, spinal muscular atrophy and cerebellar degeneration), myelodysplastic syndromes, aplastic anemia, ischemic injury associated with myocardial infarctions, stroke and reperfusion injury, arrhythmia, atherosclerosis, toxin-induced or alcohol related liver diseases, hematological diseases (including but not limited to chronic anemia and aplastic anemia), degenerative diseases of the musculoskeletal system (including but not limited to osteoporosis and arthritis) aspirin-sensitive rhinosinusitis, cystic fibrosis, multiple sclerosis, kidney diseases and cancer pain.

Compounds of formula I, as inhibitors of the cdks, can modulate the level of cellular RNA and DNA synthesis. These agents would therefore

be useful in the treatment of viral infections (including but not limited to HIV, human papilloma virus, herpesvirus, poxvirus, Epstein-Barr virus, Sindbis virus and adenovirus).

Compounds of formula I may also be useful in the chemoprevention of cancer. Chemoprevention is defined as inhibiting the development of invasive cancer by either blocking the initiating mutagenic event or by blocking the progression of pre-malignant cells that have already suffered an insult or inhibiting tumor relapse.

Compounds of formula I may also be useful in inhibiting tumor angiogenesis and metastasis.

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Compounds of formula I may also act as inhibitors of other protein kinases, e.g., protein kinase C, her2, raf I, MEK1, MAP kinase, EGF receptor, PDGF receptor, IGF receptor, PI3 kinase, wee1 kinase, Src, Abl and thus be effective in the treatment of diseases associated with other protein kinases.

The compounds of this invention may also be useful in combination (administered together or sequentially) with known anticancer treatments such as radiation therapy or with cytostatic or cytotoxic agents, such as for example, but not limited to, DNA interactive agents, such as cisplatin or doxorubicin; topoisomerase II inhibitors, such as etoposide; topoisomerase I inhibitors such as CPT-11 or topotecan; tubulin interacting agents, such as paclitaxel, docetaxel or the epothilones; hormonal agents, such as tamoxifen; thymidilate synthase inhibitors, such as 5-fluorouracil; and anti-metabolites, such as methoxtrexate.

25 Compounds of formula I may also be useful in combination with modulators of p53 transactivation.

If formulated as a fixed dose, such combination products employ the compounds of this invention within the dosage range described below and the other pharmaceutically active agent or treatment within its approved dosage range. For example, the cdc2 inhibitor olomucine has been found to act synergistically with known cytotoxic agents in inducing

apoptosis (J. Cell Sci., 108, 2897 (1995)). Compounds of formula I may also be administered sequentially with known anticancer or cytotoxic agents when a combination formulation is inappropriate. The invention is not limited in the sequence of administration; compounds of formula I may be administered either prior to or after administration of the known anticancer or cytotoxic agent. For example, the cytotoxic activity of the cyclin-dependent kinase inhibitor flavopiridol is affected by the sequence of administration with anticancer agents. Cancer Research, 57, 3375 (1997).

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The pharmacological properties of the compounds of this invention may be confirmed by a number of pharmacological assays. The exemplified pharmacological assays which follow have been carried out with the compounds according to the invention and their salts. The compounds of examples 1 to 14 exhibited cdc2/cyclin B1 kinase activity with IC50 values less than 50 μ M. The compounds of examples 1 to 14 exhibited cdk2/cyclin E kinase activity with IC50 values less than 50 μ M. The compounds of examples 1 to 14 exhibited cdk4/cyclin D1 kinase activity with IC50 values less than 50 μ M.

cdc2/cyclin B1 Kinase Assay

cdc2/cyclin B1 kinase activity was determined by monitoring the incorporation of ³²P into histone H1. The reaction consisted of 50 ng baculovirus expressed GST-cdc2, 75 ng baculovirus expressed GST-cyclin B1, 1 µg histone HI (Boehringer Mannheim), 0.2 mCi of ³²P g-ATP and 25 mM ATP in kinase buffer (50 mM Tris, pH 8.0, 10 mM MgCl₂, 1 mM EGTA, 0.5 mM DTT). The reaction was incubated at 30°C for 30 minutes and then stopped by the addition of cold trichloroacetic acid (TCA) to a final concentration of 15% and incubated on ice for 20 minutes. The reaction was harvested onto GF/C unifilter plates (Packard) using a Packard Filtermate Universal harvester, and the filters were counted on a Packard TopCount 96-well liquid scintillation counter (Marshak, D.R.,

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Vanderberg, M.T., Bae, Y.S., Yu, I.J., J. of Cellular Biochemistry, 45, 391-400 (1991), incorporated by reference herein).

cdk2/cyclin E Kinase Assay

cdk2/cyclin E kinase activity was determined by monitoring the incorporation of ³²P into the retinoblastoma protein. The reaction consisted of 2.5 ng baculovirus expressed GST-cdk2/cyclin E, 500 ng bacterially produced GST-retinoblastoma protein (aa 776-928), 0.2 mCi ³²P g-ATP and 25 mM ATP in kinase buffer (50 mM Hepes, pH 8.0, 10 mM MgCl₂, 5 mM EGTA, 2 mM DTT). The reaction was incubated at ³⁰°C for 30 minutes and then stopped by the addition of cold trichloroacetic acid (TCA) to a final concentration of 15% and incubated on ice for 20 minutes. The reaction was harvested onto GF/C unifilter plates (Packard) using a Packard Filtermate Universal harvester, and the filters were counted on a Packard TopCount 96-well liquid scintillation counter.

cdk 4/cyclin D1 Kinase Activity

cdk4/cyclin D1 kinase activity was determined by monitoring the incorporation of \$^{32}P\$ in to the retinoblastoma protein. The reaction consisted of 165 ng baculovirus expressed as GST-cdk4, 282 ng bacterially expressed as S-tag cyclin D1, 500 ng bacterially produced GST-retinoblastoma protein (aa 776-928), 0.2µCi ^{32}P \(\gamma ATP \) and 25 µM ATP in kinase buffer (50 mM Hepes, pH 8.0, 10 mM MgCl₂, 5 mM EGTA, 2 mM DTT). The reaction was incubated at 30°C for 1 hour and then stopped by the addition of cold trichloroacetic acid (TCA) to a final concentration of 15% and incubated on ice for 20 minutes. The reaction was harvested onto GF/C unifilter plates (Packard) using a Packard Filtermate Universal harvester, and the filters were counted on a Packard TopCount 96-well liquid scintillation counter (Coleman, K.G., Wautlet, B.S., Morissey, D, Mulheron, J.G., Sedman, S., Brinkley, P., Price, S., Wedster, K.R. (1997). Identification of CDK4 Sequences involved in cyclin D, and p16 binding. J. Biol. Chem. 272,30:18869-18874, incorporated by reference herein).

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Further subject matter of the invention also includes pharmaceuticals for use as described above including controlling cancer, inflammation and arthritis, which contain at least one compound of the formula I as defined above or at least one of its pharmacologically acceptable acid addition salts, and the use of a compound of the formula I as defined above for the preparation of a pharmaceutical having activity against proliferative diseases as described previously including against cancer, inflammation and/or arthritis.

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The following examples and preparations describe the manner and process of making and using the invention and are illustrative rather than limiting. It should be understood that there may be other embodiments which fall within the spirit and scope of the invention as defined by the claims appended hereto.

Example 1

N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

A. Preparation of 1-benzyloxycarbonylamino-2-butanol

A mixture of 1-amino-2-butanol (5.5 g, 61.8 mmol), benzyl 20 chloroformate (11.5 g, 67.6 mmol) and sodium carbonate (7.16 g, 67.7 mmol) in water (50 mL) was stirred at 0 °C for 3 h. Water (50 mL) was added to the reaction mixture and the product was extracted with methylene chloride (3x20 mL). The methylene chloride extract was dried over Na₂SO₄ and concentrated. The residue was passed through a short column (SiO2, hexanes: ethyl acetate /10:1; then ethyl acetate) to afford 1benzyloxycarbonylamino-2-butanol (13.9 g, 100%) as a liquid.

 1 H NMR (CDCl₃) δ 7.30 (m, 5 H), 5.45 (s, 1 H), 5.06 (s, 2 H), 3.57 (s, 1 H), 3.31 (m, 1 H), 3.04 (m, 1 H), 2.91 (m, 1 H), 1.43 (m, 2 H), 0.91 (t, J = 7.6 Hz, 3 H).

5 B. Preparation of 1-benzyloxycarbonylamino-2-butanone

To methylene chloride (60 mL) at -78 °C under argon was added oxalyl chloride (37 mL of 2 M solution in methylene chloride, 74 mmol), followed by DMSO (7.8 g, 100 mmol). The mixture was stirred at -78 °C for 20 min. and to this mixture was added a solution of 1-benzyloxycarbonylamino-2-butanol (13.9 g, 61.8 mmol) in methylene chloride (40 mL). The mixture was stirred at -78 °C for 1 h and triethylamine (21 mL) was added to the mixture. It was warmed to room temperature (rt) and washed successively with 1 N hydrochloric acid and aqueous sodium bicarbonate solution. The methylene chloride solution was dried over MgSO₄ and concentrated to afford 1-benzyloxycarbonylamino-2-butanone (11.2 g, 82%) as a solid, which was enough pure for the next reaction.

¹H NMR (CDCl₃) δ 7.32 (m, 5 H), 5.50 (s, 1 H), 5.06 (s, 2 H), 4.07 (s, 2 H), 2.43 (q, J = 7.6 Hz, 2 H), 1.06 (t, J = 7.6 Hz, 3 H).

20 C. Preparation of 1-amino-2-butanone

A solution of 1-benzyloxycarbonylamino-2-butanone (9.30 mg, 42 mmol) in ethanol (50 mL) and 1 N hydrochloric acid (46 mL) was stirred under hydrogen atmosphere in the presence of Pd/C (1.5 g, 10%) at rt for 4 h. The mixture was filtered through a celite bed and the filtrate solution was concentrated. The residue was triturated with ethyl ether to afford 1-amino-2-butanone (5.3 g, 102%) as a hydrochloride salt.

 $^{1}\rm{H}$ NMR (CD_{3}OD) δ 3.97 (s, 2 H), 2.60 (q, J = 7.6 Hz, 2 H), 1.08 (t, J = 7.6 Hz, 3 H).

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D. Preparation of 2-amino-5-thiocyanatothiazole

2-Aminothiazole (41g, 410 mM) and sodium thiocyanate (60 g, 740 mM, dried in a vacuum oven at 130 °C overnight) was dissolved in 450 mL of anhydrous methanol and the solution was cooled in a cold water bath.

5 Here was added bromine (23 mL, 445 mM) dropwise with good stirring. After the addition it was stirred for 4 h at rt. To the mixture 500 mL of water was added and it was stirred for 5 minutes, filtered through a celite bed and washed the bed with water. The pH of the filtrate solution was about 1. Most of the methanol was removed under the reduced pressure and pH of the solution was adjusted to about 7 by adding aq. sodium carbonate slowly with stirring. The precipitated solid was filtered and washed with water to obtain 37 g (57%) of the dark brown colored desired product after drying, mp 140-143 °C.

 1 H NMR (CD₃OD) δ 7.33 (s, 1H); MS (CI/NH₃) m/e 179 (M+Na)⁺, 158(M+H)⁺.

E. Preparation of of 2-acetylamino-5-thiocyanatothiazole

To a mixture of 2-amino-5-thiocyanatothiazole (15.7 g, 0.1 mol) and pyridine (12 g, 0.15 mol) in methylene chloride (100 mL) was added acetic anhydride (1.2 g, 0.12 mol) at rt. The mixture was stirred at rt for 6 h. The mixture was concentrated to dryness and to the residue MeOH (50 mL) was added. The precipitates were collected and washed with water. The solid was dried and recrystallized from MeOH to afford 2-acetylamino-5-thiocyanatothiazole (15.2 g, 76%) as a solid, mp 212 °C.

 1 H NMR (CD₃OD) δ 7.79 (s, 1H), 2.23 (s, 3 H).

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F. Preparation of [[2-(acetylamino)-5-thiazolyl]thio]acetic acid 1,1-dimethylethyl ester

To a mixture of 2-acetamino-5-thiocyanatothiazole (5.97 g, 30 mmol) in MeOH (360 mL) under argon was added dithiothreitol (9.26 g, 60

mmol) at rt. The mixture was stirred at rt for 2 h and it was concentrated to afford a reduced solid product. This solid product was dissolved in DMF (30 mL) and to this solution were added tert-butyl bromoacetate (5.85 g, 30 mmol) and potassium carbonate (5.0 g, 36 mmol). The mixture was stirred at rt for 2 h and water (200 mL) was added to the mixture. The precipitates were collected, washed with water and dried. The solid was dissolved in methylene chloride (100 mL) and MeOH (10 mL) and filtered through a silica gel pad. The filtrate solution was concentrated to afford the desired product (7.5 g, 87%) as a solid, mp 162-163 °C.

10 ¹H NMR (CDCl₃) δ 12.2 (s, 1 H), 7.48 (s, 1 H), 3.37 (s, 2 H), 2.32 (s, 3 H), 1.45 (s, 9 H); MS m/e 289 (M+H)⁺, 287 (M-H)⁻.

HPLC (Column: YMC S3 ODS 4.6x150mm; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10% MeOH-90% water-0.2% H₃PO₄; Solvent B: 90% MeOH-10% Water-0.2% H₃PO₄; UV: 220 nm): retention time 6.44 min.

G. Preparation of [[2-(acetylamino)-5-thiazolyl]thio]acetic acid

A solution of [[2-(acetylamino)-5-thiazolyl]thio]acetic acid 1,1-dimethylethyl ester (4.32 g, 15 mmol) in methylene chloride (30 mL) and trifluoroacetic acid (20 mL) was stirred at rt overnight and concentrated in vacuo. To the residue was added ethyl ether (50 mL). The precipitated solid was collected, washed with ethyl ether and dried to afford the desired product (3.38 g, 97%) as a solid, mp 210 °C.

25 ¹H NMR (CD₃OD) δ 7.48 (s, 1 H), 3.47 (s, 2 H), 2.20 (s, 3 H) ppm; MS m/e
231(M-H)⁻; HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10% MeOH-90% water-0.2%H₃PO₄; Solvent B: 90% MeOH-10% Water-0.2% H₃PO₄; UV: 254 nm): retention time 4.32 min.

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H. Preparation of [[2-(acetylamino)-5-thiazolyl]thio]-N-(2-oxobutyl)acetamide

A mixture of [[2-(acetylamino)-5-thiazolyl]thio]acetic acid (9.0 g, 38.8 mmol), HOBT (5.94 g, 38.8 mmol) and

- triethylaminopropylcarbodiimide hydrochloride salt (11.16 g, 58.2 mmol) in DMF (50 mL) was stirred at 0 °C for 0.5 h. To this mixture was added 1-amino-2-butanone hydrochloride (5.27 g, 42.7 mmol) followed by triethylamine (15 mL, 107.5 mmol). The mixture was stirred at 0 °C for 0.5 h and at rt for 1 h. Water (200 mL) was added to the mixture and the product was extracted with methylene chloride containing 10% MeOH (5x100 mL). The methylene chloride extract was dried over Na₂SO₄ and concentrated. The residue was triturated with water and the precipitated solid product was collected by filtration. It was dried to obtain the desired product (10.5 g, 90%), mp 195-196 °C.
- ¹H NMR (CDCl₃) δ 7.53 (s, 1 H), 4.14 (s, 2 H), 3.46 (s, 2 H), 2.50 (q, J = 7.6 Hz, 2 H), 2.25 (s, 3 H), 1.12 (t, J = 7.6 Hz, 3 H); MS m/e 302 (M+H)⁺. HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10%MeOH-90%water-0.2%H₃PO₄; Solvent B: 90%MeOH-10%Water-0.2%H₃PO₄; UV: 254 nm):
 retention time 4.36 min.

I. Preparation of N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

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To a solution of [[2-(acetylamino)-5-thiazolyl]thio]-N-(2-oxobutyl)acetamide (10.5 g, 34.8 mmol) in acetic anhydride (100 mL) was added conc. sulfuric acid (10 mL). The mixture was stirred at 55-60 °C for 2 h and sodium acetate (15 g, 0.18 mol) was added to the mixture. The mixture was concentrated in vacuo. To the residue was added cold water (100 mL). The precipitated solid was collected, washed with water and dried. It was purified by a flash column chromatography (SiO₂; methylene chloride:

MeOH / 100 : 5) to afford N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide (4.2 g, 43%) as a solid, mp 147-148 °C. ¹H NMR (CDCl₃) δ 12.47 (s, 1 H), 7.29 (s, 1 H), 6.61 (s, 1 H), 3.91 (s, 2 H), 2.64 (q, J = 7.6 Hz, 2 H), 2.25 (s, 3 H), 1.21 (t, J = 7.6 Hz, 3 H) ppm; MS m/e

5 284 (M+H)+;

HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10%MeOH-90%water-0.2%H₃PO₄; Solvent B: 90%MeOH-10%Water-0.2%H₃PO₄; UV: 254 nm): retention time 6.50 min.

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Example 2

N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl] benzamide

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A. Preparation of 2-amino-5-[[(5-ethyl-2-oxazolyl)methyl]thio]-thiazole

A solution of N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide (1.3 g, 4.6 mmol) in 1 N hydrochloric acid (15 mL) was stirred at 80-90 °C for 3 h. It was cooled to rt and the pH of the solution was adjusted to 7 with sodium carbonate. The product was extracted with methylene chloride (3x10 mL). The combined extract was dried over Na₂SO₄ and concentrated. The residue was triturated with ethyl ether and the precipitated solid was collected to afford 2-amino-5-[[(5-ethyl-2-oxazolyl)methyl]thio]-thiazole (610 mg, 55%) as a solid, mp 119-120 °C. 1 H NMR (CDCl₃) δ 6.93 (s, 1 H), 6.61 (s, 1 H), 5.41 (s, 2 H), 3.82 (s, 3 H), 2.62 (q, J = 7.6 Hz, 2 H), 1.18 (t, J = 7.6 Hz, 3 H); MS m/e 242 (M+H)⁺;

HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10%MeOH-90%water-0.2%H₃PO₄; Solvent B: 90%MeOH-10%Water-0.2%H₃PO₄; UV: 254 nm): retention time 3.96 min.

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B. Preparation of N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]benzamide

A mixture of 2-amino-5-[[(5-ethyl-2-oxazolyl)methyl]thio]-thiazole (48.2 mg, 0.2 mmol), benzoyl chloride (24.4 mg, 0.21 mmol) and triethylamine (35 mg, 0.35 mmol) in methylene chloride (0.5 mL) was stirred at rt for 10 min. The organic solution was washed with water and concentrated. The residue was purified by a flash column (SiO₂; hexanes: ethyl acetate / 2: 1) to afford N-[5-[[(5-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]benzamide (41 mg, 59%) as a solid, mp 122-123 °C.

1H NMR (CDCl₃) δ 12.65 (s, 1 H), 7.96 (m, 2 H), 7.61 (m,, 1 H), 7.49 (m, 2 H), 6.88 (s, 1 H), 6.56 (s, 1 H), 3.93 (s, 2 H), 2.61 (q, J = 7.6 Hz, 2 H), 1.20 (t, J = 7.6 Hz, 3 H); MS m/e 346 (M+H)⁺;
HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 mL/min; solvent system: 0-100% B in 8 min. Solvent A: 10%MeOH-90%water-

20 0.2%H₃PO₄; Solvent B: 90%MeOH-10%Water-0.2%H₃PO₄; UV: 254 nm): retention time 7.94 min.

Example 3

25 N-[5-[[(4,5-dimethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

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A. Preparation of 2-(bromomethyl)-4,5-dimethyloxazole

A mixture of 2,4,5-trimethyloxazole (0.50 mL, 4.3 mmol), N-bromosuccinimide (0.77 g, 4.3 mmol) and benzoyl peroxide (0.21 g, 0.86 mmol) in carbon tetrachloride (4 mL) was heated at 76° C under nitrogen atm.for 3 hrs. After cooling to rt, the solid was removed by filtration. The filtrate solution was washed with saturated aqueous NaHCO₃ (20 mL) and concentrated. The residue was purified by flash column chromatography (SiO₂; hexanes:ethyl acetate / 4:1) to afford 2-(bromomethyl)-4,5-dimethyloxazole (64 mg) as an yellow oil.

 $^{1}\text{H NMR (CDCl}_{3}) \delta 4.4 \text{ (s, 2 H), 2.25 (s, 3 H), 2.05 (s, 3 H).}$

B. Preparation of N-[5-[[(4,5-dimethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

N-[5-(Acetylthio)-2-thiazolyl]acetamide (0.050 g, 0.23 mmol) was dissolved in dry THF (10 ml) and here potassium*tert*-butoxide (1.0 M solution in THF, 0.25 ml, 0.25 mmol) was added to the mixture. The reaction mixture was stirred at rt for 15 min., and 2-(bromomethyl)-4,5-dimethyloxazole (0.064 g, 0.34 mmol) was added to this mixture. The reaction mixture was stirred at rt for 3 h and saturated aqueous NaHCO₃ solution (20 mL) was added to the mixture. The organic layer was separated and the aqueous layer was extracted with dichloromethane (3 x 20 mL). The combined organic layers was concentrated. The residue was purified by flash column chromatography (SiO₂; methanol:dichloromethane /1:20) to afford N-[5-[[(4,5-dimethyl-2-oxazolyl)methyl]thiol-2-thiazolyl]acetamide (15 mg, 23%) as a yellow solid. ¹H NMR (CDCl₃) δ 11.78 (s, 1 H), 7.38 (s, 1 H), 3.90 (s, 2 H), 2.30 (s, 3H), 2.22 (s 3H), 2.05 (s, 3H); MS m/e 284 (M+H)⁺; HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 ml/min; solvent system: 0-100% B in 8 min. Solvent A: 10% CH₃OH/90% H₂O/0.2%

H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 254 nm): retention time 5.87 min.

Example 4

5 N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

10 A. Preparation of diazomethane

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To a mixture of 15 ml of 40% aqueous KOH solution and 50 mL of diethyl ether at 0 °C was added 5 g (68 mmol) of N-methyl-N'-nitro-N-nitrosoguanidine in portions with stirring . The resulting mixture was stirred at 0 °C for 0.5 h. The organic phase was decanted into a dry flask and dried over solid KOH pellets to give 50 mL of diazomethane solution (ca 0.5 M, by titrating with acetic acid).

B. Preparation of 1-diazo-3,3-dimethyl-2-butanone

To the diazomethane solution at 0 °C was added a solution of 1.23

20 mL (1.21 g, 10 mmol, Aldrich) of trimethylacetyl chloride in 1 mL of diethyl ether dropwise with stirring. The resulting mixture was kept at 0 °C for 16 h. The solution was sparged with argon to remove the excess diazomethane and diethyl ether was removed under reduced pressure to give 1.33 g (10 mmol, 100%) of crude 1-diazo-3,3-dimethyl-2-butanone as a yellow liquid.

C. Preparation of 2-chloromethyl-5-t-butyloxazole

To a solution of 2 mL (2.3 g, 16 mmol) of boron trifluoride etherate in 20 mL of chloroacetonitrile at 0 °C was added a solution of 1.33 g (10

mmol) of 1-diazo-3,3-dimethyl-2-butanone in 5 mL of chloroacetonitrile dropwise. The resulting solution was stirred at 0 °C for 0.5 h. The reaction mixture was added to saturated aqueous sodium bicarbonate solution to neutralize the acid and the product was extracted three times with dichloromethane. The combined extracts was dried (sodium sulfate), concentrated and purified by flash column chromatography (Merck silica, 25 x 200 mm, dichloromethane) to give 1.1 g of 2-(chloromethyl)-5-t-butyloxazole as a yellow liquid (6.4 mmol, 64% overall from the acid chloride).

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10 ¹H NMR δ (CDCl₃): 1.30 (s, 9H), 4.58 (s, 2H), 6.68 (s, 1H); MS 174 (M+H)⁺; TLC: R_f (silica gel, dichloromethane)=0.33;
HPLC: t_R (YMC S-3 ODS 4.6x50mm rapid resolution; 2.5 ml/min, gradient 0-100% B over 8 min, Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 254 nm)= 6.5 min.

D. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

To a solution of 50 mg (0.23 mmol, Applied Chemical Laboratory) of N-[5-(acetylthio)-2-thiazolyl]acetamide in 10 mL of THF was added 0.25 mL of potassium tert-butoxide solution (1 M solution, 0.25 mmol) at rt under argon. The resulting suspension was stirred for 15 min at rt, then a solution of 59 mg of 2-(chhloromethyl)-5-t-butyloxazole (0.34 mmol) in 1 mL of THF was added. The resulting mixture was stirred at rt for 16 h, concentrated under reduced pressure and purified by flash column chromatography (silica gel, 25 x 200 mm, 1:1 EtOAc/hexanes followed by 100% EtOAc) to give 44 mg (0.14 mmol, 61%) of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide as a white solid.

¹H NMR δ (CDCl₃) 1.27 (s, 9H), 2.27 (s, 3H), 3.95 (s, 2H), 6.59 (s, 1H), 7.31 (s, 1H), 11.03 (broad s, 1H); MS 312 (M+H)⁺;

TLC: R_f (silica gel, ethyl acetate)=0.53, UV;

HPLC: retention time (YMC S-3 ODS 4.6x50mm rapid resolution; 2.5 ml/min, gradient 0-100%B over 8 min, Solvent A: 10% CH₃OH/90%

H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 254 nm)= 6.8 min.

Example 5

N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]

10 trimethylacetamide

A. Preparation of N-[(5-thiocyanato)-2-thiazolyl] trifluoroacetamide (XVIII)

To a mixture of 5-thiocyanato-2-aminothiazole (30 mmol) and 2,6-lutidine (35 mmol) in tetrahydrofuran (25 mL) and dichloromethane (50 mL) at -78 °C under argon was slowly added trifluoroaceticanhydride (33 mmol). After addition, the mixture was allowed to warm up to rt and stirred overnight. The mixture was diluted with dichloromethane (100 mL), and the organic solution was washed with 5% aqueous citric acid followed by brine, dried over magnesium sulfate and passed through a pad of silica gel. The product containing eluent was concentrated to afford 5.3 g of light brown solid.

 ^{1}H -NMR (CDCl₃) δ 12.4 (br, 1H), 7.83 (s, 1H).

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B. Preparation of 4-hydroxymethyl-3-methoxyphenyloxy Merrifield resin (XVI)

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To the suspension of sodium hydride (11.7 g, 60% in mineral oil, 293 mmol) in dimethylformamide (30 mL) at 0 °C under argon was slowly added a solution of 4-hydroxy-3-methoxybenzyldehyde (44.5 g, 292.5 mmol) in dimethylformamide (100 mL). To the resulting mixture Merrifield resin (1% DVB, from Advanced Chemtech, loading 1.24 mmol/g, 50 g, 62 mmol) and catalytic amount of tetra-n-butylammonium idodide were added, and it was heated at 65 °C for a day. The resin was filtered, washed with water (2x), 50% dimethylformamide in water (3x), dimethylformamide (2x), and methanol (5x), and dried in vacuo. The dried resin (15 g) was treated with sodium borohydride (3.4 g, 90 mmol) in tetrahydrofuran (50 mL) and ehthanol (50 mL) overnight. The resin was filtered, washed with 50% dimethylformamide in water (3x), dimethylformamide (2x), methanol (2x), and dichloromethane (5x), and dried in vacuo.

C. Preparation of 4-chloromethyl-3-methoxyphenyloxy Merrifield resin (XVII)

To a solution of triphenylphosphine (17 g, 65 mmol) in dichloromethane (200 mL) at 0 °C was slowly added triphosgene (9.2 g, 31 mmol) portionwise over a period of 30 minutes. After addition, the reaction mixture was stirred at 0 °C for 10 minutes. The solvent was removed in vacuo and the residue was redissolved in dichloromethane (200 mL). To this mixture was added 4-hydroxymethyl-3-methoxyphenyloxy Merrifield resin (12 g). The resulting mixture was agitated for 4 h. The resin was washed with dry dichloromethane (6x) and dried in vacuo.

D. Preparation of 4-[N-[(5-thiocyanato)-2-thiazolyltrifluoroacetamido]methyl]-3-methoxyphenyloxy

Merrifield resin (XIX)

A mixture of 4-chloromethyl-3-methoxyphenyloxy Merrifield resin (15g), N-[(5-thiocyanato)-2-thiazolyl]trifluoroacetamide (14 g, 55.3 mmol) and diisopropylethylamine (7.8 mL, 45 mmol) in dimethylformamide (50 mL) and dichloromethane (100 mL) was agitated overnight. The resin was washed with dimethylformamide (2x), methanol (2x), dichloromethane (4x), and dried in vacuo.

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E. Preparation of 4-[[N-[(5-mercapto)-2-thiazolyl] trifluoroacetamido]methyl]-3-methoxyphenyloxy Merrifield resin (XX)

A mixture of 4-[N-[(5-thiocyanato)-2-thiazolyltrifluoroacetamido]

methyl]-3-methoxyphenyloxy Merrifield resin (XIX, 18.5 g) and
dithiothreitol (12 g, 78 mmol) in tetrahydrofuran (100 mL) and methanol
(100 mL) was agitated overnight. The resin was washed with
dimethylformamide (2x), methanol (2x), dichloromethane (4x), and dried
in vacuo and stored under argon at -20 °C.

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F. Preparation of 4-N-[5-[[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]trifluoroacetamido]methyl-3-methoxyphenyloxy

Merrifield resin (XXI)

A stream of argon was bubbled through a mixture 4-[[N-[(5-25 Mercapto)-2-thiazolyl]trifluoroacetamido]methyl]-3-methoxyphenyloxy Merrifield resin (XX, 500 mg), halide (2.0 mmol) and 1,8-diazabicyclo[5,4,0]undec-7-ene (DBU, 1.5 mmol) in dimethylformamide (3 mL) for 5 min., and the mixture was heated at 80 °C for 2 h. The resin was washed with dimethylformamide (2x), methanol (2x), dichloromethane (4x), and dried in vacuo.

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G. Preparation of 4-N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]methyl-3-methoxyphenyloxy Merrifield resin (XXII)

A mixture of 4-N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]trifluoroacetamido]methyl-3-methoxyphenyloxy Merrifield resin (XXI, 500 mg) and sodium borohydride (4 mmol) in tetrahydrofuran (2 mL) and ethanol (2 mL) was agitated overnight. The resin was washed with 50% dimethylformamide in water (2x), dimethylformamide (2x), methanol (2x), dichloromethane (4x), and dried in vacuo.

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H. Preparation of 4-N-[5-[[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]trimethylacetamido]methyl-3-methoxyphenyloxy

Merrifield resin (XXIII)

A mixture of 4-N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2thiazolyl]methyl-3-methoxyphenyloxy Merrifield resin (XXII, 100 mg),
diisopropylethylamine (1.2 mmol) and trimethylacetyl chloride (1 mmol)
in dichloromethane (2 mL) in a polypropylene tube fitted with a
polyethylene frit and a luer stopcock was agitated overnight. The resin
was washed with dimethylformamide (2x), methanol (2x),
dichloromethane (4x), and used in the next step without drying.

- I. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]trimethylacetamide
- 4-N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2thiazolyl]trimethylacetamidolmethyl-3-methoxyphenyloxy Merrifield resin
 (XXIII) was treated with 60% trifluoroacetic acid in dichloromethane (2
 mL) in a polypropylene tube fitted with a polyethylene frit and a luer
 stopcock for 4 hours. The solution was decanted to a tube and the resin
 was washed with dichloromethane. The combined organic solution was
 concentrated in Speed Vac. The residue was purified by preparative-HPLC
 to afford 11.3 mg of the desired product.

MS m/e 354 (M+H)+.

Example 6

N-[5-[[(4-Ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

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A. Preparation of 2-(2-chloroacetamido)-1-butanol

To a mixture of 2-amino-1-butanol (5.0 mL, 53 mmol) and triethyl amine (15.0 mL, 111 mmol) in dichloromethane (20 mL) at -70 °C was added chloroacetyl chloride (4.6 mL, 58 mmol) dropwise. The reaction mixture was stirred at -70 °C for 15 min. and then was allowed to warm to rt. It was diluted with EtOAc (50 mL) and the reaction was quenched by adding water (50 mL). The organic layer was separated and the aqueous layer was extracted with EtOAc (3 x 30 mL). The combined organic layers was concentrated to afford 2-(2-chloroacetamido)-1-butanol (8.6 g, 98%) as a brown solid.

¹H NMR (CDCl₃) δ 6.75 (bs, 1 H), 4.10 (s, 2 H), 4.08(dd, 1H), 3.90 (m, 1 H), 3.68 (m, 2H), 2.98(bs, 1H), 1.60(m, 2H), 0.97 (t, 3H).

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B. Preparation of 2-(2-chloroacetamido)-1-butyraldehyde

To a solution of oxalyl chloride (14.5 mL, 29.0 mmol) in dichrolomethane (30 mL) at -78 °C DMSO (2.75 mL, 38.8 mmol) was added dropwise over 5 min.. After stirring for 10 min. at -78 °C, here was added a solution of 2-(2-chloroacetamido)-1-butanol (4.0 g, 24 mmol) in 20 mL of dichrolomethane dropwise over 15 min. The reaction mixture was stirred for 40 min. at -78 °C and here was added triethyl amine (9.4 mL, 68 mmol) dropwise over 5 min. and the reaction mixture was allowed to warm to room temperature and stirred for 2 hrs. The solid was removed by filtration and

washed with EtOAc. The organic phase was washed with 1N HCl (2 x 100 mL), saturated aqueous NaHCO₃ (1 x 10 mL) and concentrated to afford 2-(2-chloroacetamido)-1-butyraldehyde (3.7 g, 95%) as a brown oil. 1H NMR (CDCl₃) δ 9.60 (s, 1 H), 4.52 (q, 1 H), 4.12(s, 2H), 2.05 (m, 1 H), 1.80 (m, 1H), 0.97 (t, 3H).

C. Preparation of 2-chloromethy-4-ethyloxazole

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To a solution of 2-(2-chloroacetamido)-1-butyraldehyde (3.7 g, 23 mmol) in toluene (10 mL) was added POCl₃ (6.3 mL, 68 mmol). The reaction mixture was heated at 90 °C for 1 h under nitrogen. After cooling the reaction mixture to room temperature it was poured into ice water (10 mL) and the pH of the solution was adjusted to 7 with 5N NaOH. The toluene layer was separated and the aqueous layer was washed with dichloromethane (3 x 20 mL). The combined organic solution was concentrated and distilled to afford 2-chloromethy-4-ethyloxazole (1.1g, 31%) as a colorless liquid.

 $^{1}\text{H NMR (CDCl}_{3})$ δ 7.30 (s, 1H), 4.22 (s, 2 H), 2.50 (q, 2 H), 1.22 (t, 3H).

20 D. Preparation of N-[5-[[(4-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide

To a solution of 2-acetylamino-5-thiazolylthiol (0.010 g, 0.050 mmol) in dry THF (5 mL) was added potassium tert-butoxide (1.0 M solution in THF, 0.060 mL, 0.060 mmol). The reaction mixture was stirred at room temperature for 15 min. and here was added 2-chloromethyl-4-ethyloxazole (0.015 g, 0.10 mmol). After 3 h, saturated aqueous NaHCO3 solution (5 mL) was added to the mixture. The organic layer was separated and the aqueous layer was washed with dichloromethane (3 x 10 mL). The combined organic layers was concentrated. The residue was purified by flash chromatography (SiO2; methanol:dichloromethane /1:20)

to afford N-[5-[[(4-ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide (5 mg, 36%) as a white solid.

¹H NMR (CDCl₃) δ 11.25 (s, 1 H), 7.34 (s, 1 H), 7.31(s, 1H), 3.95 (s, 2 H), 2.50 (q, 2H), 2.27(s, 3H), 1.19 (t, 3H); MS m/e 284 (M+H)⁺; HPLC (Column: Zorbax Rapid resolution C-18; flow rate: 2.5 ml/min; solvent system: 0-100% B in 8 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 254 nm): retention time 6.14 min.

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Example 7

Preparation of N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-N'-cyano-N"-(2,6-difluorophenyl)guanidine.

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A solution of 100 mg of N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-aminothiazole and 68 mg of 2,6-difluorophenyl isothiocyanate was heated at 65°C for 16 hours under argon. The solution was evaporated to dryness and the residue purified by flash chromatography to give 91 mg of the intermediate thiourea.

To a solution of 30 mg of N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-N"-(2,6-difluorophenyl)thiourea, 52 mg of ethyl-3(3-

dimethylamino)propyl carbodiimide hydrochloride and 48 µL of
diisopropylethylamine in 0.5 mL methylene chloride was added a solution
of 29 mg of cyanamide in 0.1 mL tetrahydrofuran. After stirring for 1 hr,

the solvent was removed and the crude material purified by HPLC to give 8 mg of Example 636 compound.

MS: (M+H)+ 449+

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¹H NMR (400 MHz, CDCl₃): d 1.27 (9H, s), 4.19 (2H, s), 6.69 (1H, s), 7.03 (2H, m), 7.35 (1H, m), 8.74 (1H, s).

Example 8

Preparation of N-[5-[[(5-isopropyl-2-oxazolyl)fluoromethyl]thio]-2-thiazolyl acetamide.

To a stirred mixture of 2-acetamido-5-thiazole thiol acetate (141 mg) in 3 mL of dry THF under argon was added 1N t-BuOK in THF (0.72 mL). This mixture was stirred at room temperature for 25 min, and a solution of 5-isopropyl-(2-(chlorofluoromethyl))oxazole (116 mg) in 2 mL of dry THF was added. The reaction mixture was stirred at 60° C for 18 hr, diluted with 150 mL of EtOAc and washed with saturated NH₄Cl solution (2x25 mL), saturated NaHCO₃ solution (1x25 mL) and brine (1x25 mL). The organic layer was dried (MgSO₄), filtered and concentrated *in vacuo* to give Example 637 compound.

MS: (M+H)+316

HPLC retention time 3.52 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

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Example 9

Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]aminophenyl-4-(2-hydroxyethyl)sulfonamide

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A. Preparation of 5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-bromo thiazole.

To a solution of CuBr₂ (5.14 g in acetonitrile (100 mL) at 0° C was added tBuONO (4 mL, 1.2 eq) followed by 5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]amine (5.2 g). The mixture was stirred at 0° C for one hour, then at room temperature for one hour, ethyl acetate was added and the organic mixture washed with hydrochloric acid (2 X 50 mL), dried over magnesium sulfate, filtered through a pad of silica gel, and concentrated in vacuo. The residue was chromatographed on silica gel to give the bromide as an orange oil (3.9 g).

MS: (M+H) + 334

HPLC retention time 4.04 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90%

H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

B. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]aminophenyl-4-(2-hydroxyethyl)sulfonamide
A mixture of the 2-bromothiazole from Part A (0.85 g) in dimethyl
acetamide (4 mL) and 4-aminophenyl-N-(2-hydroxyethyl)sulfonamide (2.5 g, 5 eq) was stirred at 145° C for 6 hours, cooled and ethyl acetate (80 mL)
was added. The reaction mixture was washed with water (2 X 20 mL), the
combined aqueous solution was extracted with ethyl acetate, and the
combined organic layers dried over sodium sulfate, evaporated in vacuo,
and the residue was chromatographed on silica gel, then purified by
reverse phase chromatography to give N-[5-[[(5-t-butyl-2-oxazolyl)
methyl]thio]-2-thiazolyl]aminophenyl-4-(2-hydroxyethyl)sulfonamide as a
yellow solid (0.61 g).

MS: (M+H)+469

HPLC retention time 3.80 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

Example 10

Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]aminophenyl-4-sulfonamide

A mixture of the 2-bromothiazole from Example 9, Part A (106 mg) in

dimethyl acetamide (0.5 mL) and 4-aminobenzenesulfonamide (275 mg, 5

eq) was stirred at 140° C for 6 hours, cooled and the solvent was removed

under reduced pressure to provide a dark red oil which was purified by

preparative reverse phase HPLC (YMC S5 ODS) to give N-[5-[[(5-t-butyl-

2-oxazolyl)methyl]thio]-2-thiazolyl]aminophenyl-4-sulfonamide (94 mg).

MS: (M+H)+ 425

HPLC retention time 3.74 min. (Column: YMC ODS S05 4.6 X 50 mm

column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90%

15 H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220

nM).

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Example 11

20 Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-4-aminopyrimidine

To a 50 mL single necked flask was added 4-aminopyrimidine (142 mg) in

dry tetrahydrofuran (5 mL). A sodium hydride dispersion (60%, 60 mg)

25 was added, followed by heating to 60° C for one hour. The solution of the

anion was cooled to room temperature and the 2-bromothiazole from

Example 9, Part A (100 mg) was added. The reaction was heated for 24

hours at 60° C, cooled to room temparature, quenched with hydrochloric

acid and partitioned between water and ethyl acetate (25 mL each). The

organic layer was washed with water (2 X 25 mL), brine (25 mL), dried

over sodium sulfate and concentrated in vacuo to give a solid, which was

purified by trituration with 1:1 ethyl acetate:hexanes to give N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-4-aminopyrimidine (42 mg).

MS: (M+H)+ 348

HPLC retention time 3.63 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

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Example 12
Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-

thiazolyl]-3-(hydroxymethyl)aniline

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A. Preparation of N-2-[3-(hydroxymethyl)phenyl]aminothiazole

To a solution of 3-hydroxymethyl aniline (2.46 g) in dry tetrahydrofuran (50 mL) at -78° C was added methyl lithium-lithium bromide solution in ether (27 mL of 1.5 M solution). The reaction mixture was stirred at -78°

C for 10 minutes, warmed to room temperature for 10 minutes, and then cooled to -78° C and 2-bromothiazole (1.31 g) was added. The reaction mixture was stirred at 0° C for one hour, then at room temperature for 3 hours, quenched by addition of hydrochloric acid (20 mL of 2N solution), concentrated and extracted with ethyl acetate. The combined organic extracts were dried over sodium sulfate, concentrated and chromatographed on silica gel to give N-2-[3-(hydroxymethyl)phenyl] aminothiazole (0.68 g).

10 B. Preparation of N-2-[3-(hydroxymethyl)phenyl]aminothiazole-5-thiocyanate

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To a cooled solution(ice-salt bath) of the compound of part A (680 mg) and ammonium thiocyanate (500 mg) in methanol (35 mL) was added portionwise bromine (0.21 mL). After disappearance of the bromine color the reaction was concentrated and partitioned between ethyl acetate and water. The aqueous layer was extracted with ethyl acetate and the combined organic extracts were dried over sodium sulfate, concentrated and chromatographed on silica gel to give N-2-[3-(hydroxymethyl)phenyl] aminothiazole-5-thiocyanate as a yellow solid (490 mg).

C. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-3-(hydroxymethyl)aniline

To a dark red solution of the thiocyanate of part B (490 mg) in tetrahydrofuran/ethanol was added sodium borohydride portionwise (84

mg). After gas evolution had ceased, acetone (0.65 mL) was added the reaction stirred for 8 minutes, followed by addition of 2-chloromethyl-5-t-butyl-oxazole (Example 5, Part C compound, 0.5 g) and the reaction stirred for one hour at room temperature. The reaction was concentrated,

5 extracted with ethyl acetate, the combined organic extracts dried over sodium sulfate, and filtered through a pad of silica gel to provide the product (0.69 g).

MS: (M+H)+ 376

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HPLC retention time 3.84 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

Example 13

15 Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-aminopyridine

A. Preparation of N-2-[pyridinyl]aminothiazole

To a suspension of sodium hydride (60% suspension, 1.8 g) in tetrahydrofuran (200 mL) was added portionwise 2-aminopyridine (4.23 g), and the mixture was slowly heated to 55° C for 30 minutes. The reaction mixture was then cooled to -10 deg C and a solution of 2-

bromothiazole (2.46 g) in tetrahydrofuran (2 mL) was added dropwise. The reaction mixture was stirred at 55° C for 5 hours, cooled and quenched with hydrochloric acid (2N, 20 mL), concentrated, and ethyl acetate was added. The resulting solid was filtered to give N-2-[pyridinyl]aminothiazole (1.41 g).

B. Preparation of N-2-[pyridinyl]-5-bromo-aminothiazole

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To a solution of the compound of Part A(0.88 g) in acetic acid(15 mL) was added bromine (0.22 mL in 2 mL acetic acid) dropwise at room temperature. The reaction mixture was stirred at room temperature for 2 hours, the was solvent removed under reduced pressure, and the resulting solid was triturated with ether to provide N-2-[pyridinyl]-5-bromo-aminothiazole (1.6 g) as the hydrobromide salt.

C. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-aminopyridine

To a solution of N-2-[pyridinyl]-5-bromo-aminothiazole (8 g) and 2
thioacetyl-5-t-butyl oxazole (8 g) in methanol (500 mL) under argon was added a degassed solution of sodium hydroxide (25 mL of 3 N solution) at room temperature. The reaction mixture was stirred for 20 minutes and then heated to 60° C for one hour, concentrated in vacuo, partitioned between water (125 mL) and ethyl acetate (500 mL) and the aqueous layer was back-extracted (2 X 125 mL) with ethyl acetate. The combined organic layers were washed with brine (25 mL), dried over sodium

sulfate, filtered through a pad of silica gel, and the solvents removed in vacuo. The solid residue was recrystallized form ethyl acetate to provide N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-aminopyridine (7.5 g).

5 MS: (M+H)+ 347

HPLC retention time 4.01 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90% H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).

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Example 14

Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-[5-[(((3-hydroxy-2,2-dimethyl)propyl)amino)methyl]] aminopyridine

A. Preparation of N-2-[(5-bromo)pyridinyl]aminothiazole

To a suspension of sodium hydride (60% suspension, 5.2 g) in tetrahydrofuran (150 mL) was added portionwise 2-amino-4-bromopyridine (15 g), and the mixture was stirred at room temperature for 15 minutes. 2-Bromothiazole (3.8 mL) was added, and the reaction mixture was stirred at room temperature for one hour and then heated at reflux temperature for 2.5 hours, cooled, quenched with 6% citric acid and

extracted with ethyl acetate (2 X 100 mL). The organic layers were concentrated, dried over magnesium sulfate and the filtrate concentrated in vacuo to give a dark brown residue which was triturated with ether/hexanes to provide N-2-[(5-bromo)pyridinyl]aminothiazole as a yellow solid (8.9 g)

B. Preparation of N-2-[(5-carboxaldehyde)pyridinyl] aminothiazole

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A suspension of the Part A compound (6.4 g) in tetrahydrofuran (300 mL) was heated to reflux to effect dissolution, the reaction mixture was cooled to -70° C and treated with t-BuMgCl (13 mL of 2M solution in ether) dropwise over 10 minutes. The temperature was raised to -55° C, and t-BuLi (36 mL of 1.7 M solution in hexanes) was added dropwise, and the reaction mixture stirred for 20 minutes. The reaction mixture was then cooled to -70° C and DMF (8 mL) was added, the resulting mixture was stirred at -50° C for one hour and then warmed to 0° C over one hour, quenched with acetic acid (8 mL) and partitioned between ethyl acetate and water (300 mL each). The aqueous layer was back extracted with ethyl acetate (2 X 200 mL) and the combined organic layers dried over magnesium sulfate and concentrated, the solid washed with ethyl acetate and ether, and dried to give N-2-[(5-carboxaldehyde) pyridinyl] aminothiazole (3.15 g).

C. Preparation of N-2-[(5-carboxaldehyde)pyridinyl]-5-bromo-aminothiazole

A solution of N-2-[(5-carboxaldehyde) pyridinyl] aminothiazole(0.5 g) in

acetic acid (6 mL) and dichloromethane (20 mL) was treated with bromine

(0.12 mL) in dichloromethane (3 mL). The reaction mixture was stirred

for 30 minutes at room temperature, ether was added, and the resulting

precipitate was collected by filtration, washed with ether to give N-2-[(5-carboxaldehyde)pyridinyl]-5-bromo-aminothiazole (0.69 g).

D. Preparation of N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-aminopyridine-5-carboxaldehyde

10

To a solution of the compound of Part C (3.8 g) and 5-t-butyl-2-(S-

isothiourea)methyl oxazole (3.06 g) in methanol (300 mL) under nitrogen was added degassed sodium hydroxide (6.4 g of 50% w/w solution). The reaction mixture was heated at 76° C for 6 hours, the methanol was removed in vacuo, water was added, and the solid was collected by filtration, washed with water and ethyl acetate, and dried to give N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2- thiazolyl]-2-aminopyridine-5-carboxaldehyde (0.53 g). The filtrate was extracted with ethyl acetate (4 X

5.Zz=

200 mL), dried over magnesium sulfate, and concentrated in vacuo and

triturated with ether/ethyl acetate to give an additional 2.02 g of the desired compound.

- E. Preparation of N-[5-[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-[5-[((3-hydroxy-2,2-dimethyl)propyl)amino)methyl]] aminopyridine
 To a solution of the aldehyde of Part D (1.5 g) and 3-amino-2,2-dimethyl propanol (2.06 g) in tetrahydrofuran (100 mL) was added sodium triacetoxyborohydride (6.0 g), followed by acetic acid (5 mL). The reaction mixture was stirred for 30 minutes at room temperature, and the solvents removed in vacuo to give a yellow solid which was purified by column chromatography to give N-[5-[((5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-[5-[(((3-hydroxy-2,2-dimethyl)propyl)amino)methyl]] aminopyridine (1.08 g).
- MS: (M+H)+ 462
 HPLC retention time 3.22 min. (Column: YMC ODS S05 4.6 X 50 mm column, 0% to 100% B gradient in 4 min. Solvent A: 10% CH₃OH/90%
 H₂O/0.2% H₃PO₄; Solvent B: 90% CH₃OH/10% H₂O/0.2% H₃PO₄; UV: 220 nM).
- Using the procedures described herein or by modification of the procedures described herein as known to one or ordinary skill in the art, the following additional compounds have been prepared and disclosed in Table 1:

Example	Structure	Molecular Formula	(M+H)+
15	H _M S S S	C9H11N3OS2	242
16		C12H15N3O2S2	298
17	The state	C13H17N3O2S2	312
18		C11H10F3N3O2S2	338
19		C14H19N3O2S2	326
20		C21H17N3O2S2	408
21		C17H24N4O2S2	381
22		C17H17N3O2S2	360

Example	Structure	Molecular Formula	(M+H)+
23		C15H19N3O2S2	338
24		C17H17N3O3S2	376
25	J. J. J.	C17H23N3O2S2	366
. 26		C14H19N3O2S2	326
27		C13H15N3O2S2	310
28	7 September 1988	C15H13N3O2S2	332
29	١٠٠١	C13H11N3O2S2	306
30	THIS S	C10H11N3O2S2	270
31	THIS SOLD	C12H15N3O2S2	298

Example	Structure	Molecular Formula	(M+H)+
32	S S S S S S S S S S S S S S S S S S S	C13H16BrN3O2S2	391
33	J. S.	C15H12FN3O2S2	350
34		C13H15N3O4S2	342
35	4	C15 H21 N3 O2 S2	340
36	-t0	C19H21N3O2S2	388
37		C18H17N3O4S2	404
38		C15H19N3O4S2	370
39		C14H17N3O4S2	356
40	S S N	C16H19N3O3S2	366

Example	Structure	Molecular Formula	(M+H)+
41	S S NH O	C16H21N3O4S2	384
42	J. S. S. MH	C15H19N3O4S2	370
43	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	C16H21N3O4S2	384
44		C18 H17 N3 O4 S2	404
45	John Strain	C15H19N3O4S2	370
46	Chia.	C16 H14 F N3 O2 S2	364
47	Q'f-12,	C16 H14 CI N3 O2 S2	380
48	Land of the state	C16 H13 Cl2 N3 O2 S2	415
49		C18 H19 N3 O4 S2	406

Example	Structure	Molecular Formula	(M+H)+
50	-Gia	C18 H19 N3 O4 S2	406
51		C18 H19 N3 O4 S2	406
52		C18 H19 N3 O2 S2	374
53		C18 H20 N4 O2 S2	503
54		C17 H17 N3 O2 S2	360
55		C18 H19 N3 O2 S2	374
56		C18 H19 N3 O2 S2	374
57		C18 H20 N4 O2 S2	503

Example	Structure	Molecular Formula	(M+H)+
58	dia	C18 H20 N4 O2 S2	503
59		C19 H16 N4 O2 S2	511
60	d d	C18 H16 N4 O2 S2	499
61	To sell the	C18 H16 N4 O2 S2	499
62	dia g	C16 H13 F2 N3 O2 S2	382
63	print.	C17 H15 CI F N3 O2 S2	412
64		C19 H19 N3 O4 S2	418
65	The state of	C18 H16 F3 N3 O2 S2	428

Example	Structure	Molecular Formula	(M+H)+
66		C17 H16 F N3 O2 S2	378
67		C17 H16 N4 O4 S2	405
68		C17 H16 N4 O4 S2	· 405
69	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	C19 H21 N3 O4 S2	420
70		C19 H17 N3 O3 S2	400
71	LONS STANDO	C12 H15 N3 O3 S2	314
72	La stallo	C13 H17 N3 O3 S2	328
. 7 3		C15 H14 N4 O2 S2	461
, 7 4	Lange Ho	C16 H19 N3 O2 S2	350

Example	Structure	Molecular Formula	(M+H)+
75	La state	C15 H17 N5 O2 S2	364
76		C13 H14 F3 N3 O2 S2	366
77	Land Stand	C15 H15 N3 O2 S3	366
78		C17 H23 N3 O2 S2	3 6 6
79		C16 H16 N4 O2 S2	475
80	LONS SHIP NH,	C12 H16 N4 O2 S2	427
· 81		C18 H19 N3 O3 S2	-390
82	This is	C18 H18 N4 O3 S2	403
83	Cha.	C22 H19 N3 O3 S2	438

Example	Structure	Molecular Formula	(M+H)+
84		C17 H17 N3 O3 S2	376
85		C22 H19 N3 O2 S2	422
86		C16 H14 CI N3 Q2 S2	380
87	d'aris	C17 H17 N3 O3 S2	376
88		C16 H14 CI N3 O2 S2	380
89		C17 H17 N3 O3 S2	376
90		C17 H15 N3 O4 S2	390
91		C17 H14 N4 O2 S3	403

Example	Structure	Molecular Formula	(M+H)+
92		C17 H16 CI N3 O2 S2	394
93		C18 H19 N3 O3 S2	390
94	Of the	C19 H19 N3 O2 S2	386
95	TO DE	C21 H23 N3 O2 S2	414
96		C17 H16 CI N3 O2 S2	394
97	The state of the s	C18 H19 N3 O3 S2	390
98		C17 H16 CI N3 O2 S2	394
99		C18 H17 N3 O4 S2	404

Example	Structure	Molecular Formula	(M+H)+
100	of the s	C25 H22 N4 O2 S2	589
101	Ly style	C14 H17 N3 O3 S2	340
102	Ly style	C14 H17 N3 O3 S2	340
103		C15 H14 N4 O2 S2	461
104		C16 H21 N3 O2 S2	352
105	400	C18 H17 N3 O3 S2	3 <u>8</u> 8
106	T T T T T T T T T T T T T T T T T T T	C16 H16 N4 O2 S2	475
107		C19 H18 N4 O2 S2	513

Example	Structure ;	Molecular Formula	(M+H)+
108		C17 H14 N4 O2 S2	371
109		C20 H17 N3 O2 S2	396
110		C21 H18 N4 O3 S2	553
111	oning	C23 H21 N3 O3 S2	452
112	St. of	C20 H21 N3 O2 S2	400
113	of of the	C22 H23 N3 O3 S2	442
114	(A) (A)	C17 H15 N5 O2 S2	500
115	\$\frac{1}{2} \frac{1}{2} \frac	C18 H18 N4 O3 S2	403

Example	Structure	Molecular Formula	(M+H)+
116	را بر ب	C17 H17 N5 O2 S3	420
117		C17 H16 Br N3 O2 S2	439
118		C17 H16 F N3 O2 S2	378
119		C17 H15 Cl2 N3 O2 S2	429
120		C17 H15 N3 Ö3 S2	374
121		C18 H19 N3 O2 S2	374
122	The state of the s	C17 H16 Br N3 O2 S2	439
123	A CANA	C18 H19 N3 O2 S2	374

Example	Structure	Molecular Formula :	(M+H)+
124	N S H Br	C17 H16 Br N3 O2 S2	439
125		C18 H19 N3 O2 S2	374
126		C18 H16 N4 O2 S2	499
127	S. A.	C17 H15 F2 N3 O2 S2	396
128		C17 H15 F2 N3 O2 S2	396
129	So-City	C17 H15 F2 N3 O2 S2	396
130		C20 H23 N3 O2 S2	402
131		C18 H19 N3 O3 S2	390

Example	Structure	Molecular Formula	(M+H)+
132	Chiral N Chiral	C17 H18 N4 O2 S2	489
133	Long Styles	C14 H17 N3 O2 S2	324
134	Ly stipe	C13 H17 N3 O3 S2	328
135	Long Styles	C14 H13 N3 O3 S2	336
136	ST ST N	C14 H13 N3 O3 S2	336
137	La still	C15 H21 N3 O2 S2	340
138	La Sight	C15 H21 N3 O2 S2	340
139	Long Land	C15 H21 N3 O2 S2	340
140	Land Stand	C15 H21 N3 O2 S2	340
141	CHQ.	C14 H13 N5 O2 S2	348

Example	Structure	Molecular Formula	(M+H)+
142	La Santo	C15 H15 N3 O3 S2	350
143	Ly Stilo	C14 H17 N3 O4 S2	356
144		C14 H15 N5 O2 S2	464
145		C19 H21 N3 O2 S2	388
146	South Contraction of the contrac	C16 H16 N4 O2 S2	475
147	de la companya della companya della companya de la companya della	C19 H18 N4 O2 S2	513
148		C15 H17 N5 O2 S2	478
149		C19 H21 N3 O3 S2	404
150	S S H H Oran	C12 H16 N4 O2 S2	427

Example	Structure	Molecular Formula	(M+H)+
151	the state of	C20 H20 N4 O2 S2	527
152	La stall NHS	C13 H18 N4 O2 S2	441
153		C19 H18 N4 O4 S2	431 .
154	Long Linko	C14 H17 N3 O2 S2	324
155		C15 H21 N3 O2 S2	340
156	NO STATE OF THE ST	C13 H14 N4 O3 S3	371
157	La STATA	C15 H20 N4 O2 S2	467
158	La stando	C17 H22 N4 O3 S2	395
159		C14 H17 N3 O2 S2	324
160	475-63	C19 H18 N4 O2 S2	513

Example	Structure	Molecular Formula	(M+H)+
161	Comment of the Commen	C14 H19 N3 O2 S2	326
162	of the	C19 H21 N3 O2 S2	388
163	-C;+A	C16 H13 Cl2 N3 O2 S2	415
164		C17 H17 N3 O2 S2	360
165	dia	C16 H12 F3 N3 O2 S2	400
166		C20 H18 N4 O2 S2	525
167	Contraction of the second of t	C20 H18 N4 O2 S2	525
168		C19 H21 N3 O2 S2	388
169	in this	C19 H21 N3 O4 S2	420

Example	Structure	Molecular Formula	(M+H)+
170		C17 H16 F N3 O2 S2	378
171		C20 H23 N3 O5 S2	450
172		C18 H16 F3 N3 O2 S2	428
173	S, Dra	C19 H21 N3 O2 S2	388
174		C19 H21 N3 O2 S2	388
175		C18 H19 N3 O2 S2	374 -
176	Crès de la	C17 H17 N3 O3 S2	376
177		C19 H22 N4 O2 S2	517

Example	Structure	Molecular Formula	(M+H)+
178		C19 H21 N3 O2 S2	388
179	train	C19 H21 N3 O4 S2	420
180		C17 H15 F2 N3 O2 S2	396
181	La still	C14 H15 N5 O2 S2	350
182	La stall	C15 H14 N4 O2 S2	461
183	Chieal	C18 H19 N3 O3 S2	390
184	is the second of	C18 H19 N3 O4 S2	406
185		C22 H19 N3 O3 S2	438
186	The state	C17 H16 N4 O4 S2	405

Example	Structure	Molecular Formula	(M+H)+
187		C20 H23 N3 O2 S2	402
188	Jip y	C23 H21 N3 O2 S2	436
189	Shirt	C24 H23 N3 O2 S2	450
190		C23 H21 N3 O2 S2	436
191		C21 H19 N3 O2 S2	410
192		C21 H19 N3 O2 S2	410
193	in a	C17 H15 Cl2 N3 O2 S2	429
194	Forg	C19 H21 N3 O4 S2	420

Example	Structure	Molecular Formula	(M+H)+
195	Chiral Chiral	C18 H19 N3 O2 S2	374
196	Siring O	C19 H18 F3 N3 O3 S2	458
197		C22 H27 N3 O2 S2	430
198		C18 H19 N3 O2 S2	374
199	المراجل المراج	C12 H15 N3 O2 S2	298
200	J. J	C18 H26 N4 O4 S2	427
201	the state of the s	C12 H13 N3 O4 S2	328
202		C11 H13 N3 O4 S2	316
203		C11 H13 N3 O3 S2	. 300

Example	Structure	Molecular Formula	(M+H)+
204	H ₂ N S O	C11 H15 N3 O S2	270
205	HAN S S ON N	C10 H13 N3 O S2	256
206	W.	C17 H16 N4 O4 S2	405
207	cox o	C19 H20 N4 O2 S2	401
208		C16 H15 Br N4 O2 S2	440
209	in the second se	C17 H16 N6 O2 S2	515
210	w.to	C19 H17 N5 O2 S2	526
211	ooto	C20 H23 N5 O3 S2	560

Example	Structure	Molecular Formula	(M+H)+
212		C16 H16 N4 O2 S2	361
213		C16 H14 F2 N4 O2 S2	397
214		C16 H15 CI N4 O2 S2	395
215	75. CF.	C17 H18 N4 O3 S2	391
216	No. of	C17 H18 N4 O2 S2	375
217		C16 H15 Br N4 O2 S2	44 0
218		C16 H15 CI N4 O2 S2	395
219		C16 H14 Cl2 N4 O2 S2	430

Example	Structure	Molecular Formula	(M+H)+
220		C17 H17 CI N4 O3 S2	425
221	De Of	C17 H18 N4 O3 S2	391
222	di Hara	C16 H15 Br N4 O2 S2	440
223	Sisterior of the second	C16 H15 F N4 O2 S2	379
224	Janati.	C17 H18 N4 O2 S2	375
225	rough.	C17 H18 N4 O3 S2	391
226	ې نتمر	C16 H15 CI N4 O2 S2	395
227	STOY.	C18 H19 N5 O3 S2	418

Example	Structure	. Molecular Formula	(M+H)+
228	-ord	C17 H18 N4 O3 S2	391
229	pro,	C18 H21 N5 O2 S2	518
230		C16 H15 F N4 O2 S2	379
231		C16 H15 F N4 O2 S2	379
232	Derigh.	C17 H18 N4 O2 S2	375
233	Traff.	C17 H17 N5 O3 S2	4 04
234	5) 1-1-21	C17 H15 N5 O2 S3	418
235		C17 H16 N6 O2 S2	401

Example	Structure	Molecular Formula	(M+H)+
236		C16 H15 N7 O2 S2	402
237	C July 100	C16 H17 N5 O2 S2	490
238		C15 H20 N4 O2 S2	353
239	Orthis I	C17 H17 CI N4 O2 S2	409
240		C17 H19 N5 O2 S2	504
241		C17 H19 N5 O2 S2	504
242	A A A A A A A A A A A A A A A A A A A	C19 H18 N6 O2 S3	459
243	المراباء المراباء	C15 H16 N4 O2 S3	381
244	Califorate (A)	C15 H20 N4 O3 S2	369

Example	Structure	Molecular Formula	(M+H)+
245		C16 H20 N6 O2 S2	507
246		C18 H25 N5 O4 S2	440
247	225	C17 H24 N4 O2 S2	381
248		C18 H20 N4 O2 S2	389
249		C17 H18 N4 O2 S2	375
250	dyby	C18 H20 N4 O2 S2	389
251	s ait	C19 H22 N4 O2 S2	403
252		C17 H19 N5 O2 S2	504
253	A A A A A A A A A A A A A A A A A A A	C17 H17 CI N4 O2 S2	409

Example	Structure	Molecular Formula	(M+H)+
254	C. J.	C16 H17 N5 O2 S2	490
255		C17 H25 N5 O2 S2	510
256	Q to	C16 H17 N5 O2 S2	490
257	Joseph Joseph	C17 H25 N5 O2 S2	510
258		C18 H20 N4 O2 S2	389
259	المرابط المرا	C15 H16 N4 O3 S2	365
260		C17 H16 F2 N4 O2 S2	411
261		C15 H22 N4 O2 S2	355
262	مار المراج ا	C14 H18 N4 O2 S2	339
263		C14 H20 N4 O2 S2	341

Example	Structure	Molecular Formula	(M+H)+
264	٢٦٩٤٥٥	C15 H22 N4 O2 S2	355
265	J. Jing.	C17 H17 CIN4 O2 S2	409
266	D TO STATE OF THE PARTY OF THE	C18 H20 N4 O2 S2	389
267	E Starter	C18 H20 N4 O3 S2	. 405
268	4,000	C18 H20 N4 O3 S2	405
269		C18 H20 N4 O3 S2	405
270		C16 H22 N4 O3 S2	341
271	Joseph -	C14 H20 N4 O2 S2	512
272	The Color	C17 H27 N5 O2 S2	353
273		C16 H22 N4 O3 S2	425

Example	Structure	Molecular Formula	(M+H)+
274		C18 H24 N4 O4 S2	401
275		C19 H20 N4 O2 S2	383
276		C17 H26 N4 O2 S2	355
277	Joseph .	C15 H22 N4 O2 S2	433
278		C19 H20 N4 O4 S2	512
279		C16 H21 N5 O3 S2	353
280		C15 H20 N4 O3 S2	367
281		C16 H22 N4 O2 S2	389
282		C16 H21 N5 O3 S2	425
283	Tip is	C18 H24 N4 O4 S2	369

Example	Structure	Molecular Formula	(M+H)+
284	Joseph L	C13 H18 N4 O2 S2	465
285		C13 H14 N6 O2 S2	493
286	Sold of	C15 H18 N6 O2 S2	466
287	Jos State	C12 H13 N7 O2 S2	366
288	Joseph.	C14 H15 N5 O3 S2	366
289	Sold of the second	C13 H14 N6 O2 S3	409
290		C17 H17 CI N4 O2 S2	387
291	Christ.	C18 H18 N4 O2 S2	375
292		C17 H18 N4 O2 S2	405

Example	Structure	Molecular Formula	(M+H)+
293		C18 H20 N4 O3 S2	389
294	somit	C17 H16 F2 N4 O2 S2	490
295		C16 H17 N5 O2 S2	476
296	De Chi	C15 H15 N5 O2 S2	510
297	- And	C15 H14 CI N5 O2 S2	490
298	January.	C16 H17 N5 O2 S2	490
299	Jana de	C16 H17 N5 O2 S2	476
300	W.C.	C15 H15 N5 O2 S2	526

Example	Structure	Molecular Formula	(M+H)+
301		C15 H15 N5 O2 S2	540
302		C18 H29 N5 O2 S2	526
303	THYS S OF Y	C14 H19 N3 O2 S2	326
304		C21 H23 N3 O2 S2	414
305		C19 H25 N3 O2 S2	392
306	المريخ. وي	C22 H21 N3 O2 S2	424
307	w.	C22 H21 N3 O2 S2	424
308		C15 H19 N3 O2 S2	338

Example	Structure	Molecular Formula	(M+H)+
<i>)</i> 309		C16 H23 N3 O2 S2	354
310		C18 H19 N3 O2 S2	374
311		C18 H16 N4 O2 S2	385
312	Q, and	C20 H23 N3 O2 S2	402
313		C18 H17 F2 N3 O2 S2	410
314	J'Hay	C21 H23 N3 O2 S2	414: :
315		C18 H16 N4 O2 S3	417
316		C19 H19 N3 O4 S2	418

Example	Structure	Molecular Formula	(M+H)+
317	Chity in	C20 H23 N3 O3 S2	418
318		C18 H18 N4 O4 S2	419
319		C18 H18 N4 O4 S2	-419
320		C18 H18 N4 O4 S2	419
321		C19 H21 N3 O4 S2	420
322	40,40	C19 H21 N3 O4 S2	420
323	SAMA NHH	C18 H19 N5 O2 S3	434
324	d'i	C18 H19 N5 O2 S3	434

Example .	Structure	Molecular Formula	(M+H)+
32 5		C19 H18 F3 N3 O2 S2	442
326		C18 H18 Br N3 O2 S2	453
327		C21 H25 N3 O5 S2	464
328		C23 H28 N4 O4 S2	489
329		C20 H21 N3 O2 S2	400
330	0,00	C18 H25 N3 O2 S2	380
331	والمراز	C19 H21 N3 O2 S2	388
332	ard.	C27 H26 N4 O3 S2	519

Example	Structure	Molecular Formula	(M+H)+
333		C19 H21 N3 O3 S2	404
334		C20 H23 N3 O2 S2	402
335		C19 H21 N3 O2 S2	388
336		C19 H21 N3 O2 S2	388
337		C19 H21 N3 O3 S2	404
338	\$2,50°	C26 H28 N4 O4 S3	557 -
339	04	C19 H27 N3 O2 S2	394
340	5 to	C22 H22 N4 O3 S2	455

Example i	Structure	Molecular Formula	(M+H)+
341		C22 H25 N3 O4 S2	460
342	ching.	C20 H21 N3 O3 S2	416
343	فهجور	C15 H19 N3 O4 S2	370
344	Local Contraction	C20 H18 F3 N3 O2 S2	454
345	La Caria	C24 H26 N4 O3 S2	483
346		C18 H19 N3 O3 S2	390
347	Line Sell on	C18 H19 N3 O3 S2	390
348		C20 H20 N4 O2 S2	413
349	ora's	C18 H19 N3 O2 S2	374

Example	Structure	Molecular Formula	(M+H)+
350		C19 H18 N4 O2 S2	399
351		C17 H18 N4 O2 S2	489
352		C17 H18 N4 O2 S2	489
353	Stard Co	C20 H20 N4 O2 S2	413
354		C20 H24 N4 O2 S2	531
355	prais	C21 H22 N4 O2 S2	427
356		C16 H17 N5 O4 S2	408
357	\$ 150 HQ	C19 H18 N6 O2 S3	687

Example	Structure	Molecular Formula	(M+H)+
358	~ SS	C11 H15 N3 O S2	270
359		C17 H19 N3 O S2	346
360		C13 H19 N3 O S2	. 298
361	J'III	C22 H25 N3 O2 S2	428
362	+2, 2, 3, 4	C20 H27 N3 O2 S2	406
363	The contract of the contract o	C23 H23 N3 O2 S2	438
364	on of	C23 H23 N3 O2 S2	438
365	+4	C16 H21 N3 O2 S2	352
366	+9	C17 H25 N3 O2 S2	368
367		C19 H21 N3 O2 S2	388

Example	Structure	Molecular Formula	(M+H)+
368		C19 H18 N4 O2 S2	399
369	Q tot	C21 H25 N3 O2 S2	416
370	X - Ai	C19 H19 F2 N3 O2 S2	424
371		C22 H25 N3 O2 S2	428
372	Willy	C19 H18 N4 O2 S3	431
373	in the second	C20 H21 N3 O4 S2	432
374	5,57	C21 H25 N3 O3 S2	432
375	troi	C19 H20 N4 O4 S2	433

Example ·	Structure	Molecular Formula	(M+H)+
376	T. C. C.	C19 H20 N4 O4 S2	433
377		C20 H23 N3 O4 S2	434
378		C20 H23 N3 O4 S2	434
379		C19 H21 N5 O2 S3	448
380		C19 H21 N5 O2 S3	448
381		C19 H20 Br N3 O2 S2	467
382	- jara	C22 H27 N3 O5 S2	478
383	the state of the s	C24 H30 N4 O4 S2	503
384	OS Diggs	C21 H23 N3 O2 S2	414

Example	Structure	Molecular Formula	(M+H)+ ·
385	O.O.	C19 H27 N3 O2 S2	394
386	S. Jarot	C20 H23 N3 O2 S2	402
387	Stragt .	C28 H28 N4 O3 S2	533
388	W. J.	C20 H23 N3 O3 S2	418
389	Xi Vi	C19 H20 N4 O5 S2	449
390	Q, is to	C21 H25 N3 O2 S2	416
391	Birst.	C25 H27 N3 O3 S2	482
392	Quot to	C20 H23 N3 O2 S2	402

Example :	Structure 1	Molecular Formula	(M+H)+
393	Children to our	C20 H23 N3 O2 S2	402
394	China Cond	C20 H23 N3 O3 S2	418
395	Starst	C18 H20 N4 O2 S2	503
396	of the set	C27 H30 N4 O4 S3	571
397	O.C.	C20 H29 N3 O2 S2	408
398	ON SH	C23 H24 N4 O3 S2	469
. 399	group	C23 H27 N3 O4 S2	474
400	S. P. D.	C21 H23 N3 O3 S2	430

Example	Structure	Molecular Formula	(M+H)+
401	+4	C16 H21 N3 O4 S2	384
402	XI XI	C21 H20 F3 N3 O2 S2	468
403	A Constant	C25 H28 N4 O3 S2	497
404	Service.	C19 H21 N3 O3 S2	404
405		C21 H22 N4 O2 S2	427
406		C20 H20 N4 O2 S2	413
407	ord Ord	C18 H20 N4 O2 S2	503
408	ora px	C18 H20 N4 O2 S2	503

Example	Structure	Molecular Formula :	(M+H)+
409	The state of the s	C21 H22 N4 O2 S2	427
410	to gx	C21 H26 N4 O2 S2	545
411	torque	C22 H24 N4 O2 S2	441
412	train.	C16 H19 N5 O2 S3	524
413	3,47	C20 H23 N3 O3 S2	418
414	+4	C16 H19 N5 O2 S2	492
415	A.C.	C17 H19 N5 O4 S2	422
416	the first	C26 H34 N4 O4 S2	531

Example	Structure	Molecular Formula	(M+H)+
417	tions.	C24 H30 N4 O4 S2	503
418	tro	C25 H32 N4 O4 S2	517
419		C21 H26 N4 O2 S2	545
420	of of	C19 H22 N4 O2 S2	517
421	C. J.	C20 H24 N4 O2 S2	531
422	Joseph Property	C19 H22 N4 O2 S2	403
423		C16 H14 F2 N4 O2 S2	. 397
424		C16 H14 Cl2 N4 O2 S2	430

Example	Structure :	Molecular Formula	(M+H)+
425		C18 H20 N4 O S3	405
426		C16 H14 Cl2 N4 O S3	446
427		C21 H23 N3 O2 S2	414
428	Jan	C19 H25 N3 O2 S2	392
429	the state of the s	C22 H21 N3 O2 S2	424
430	aro conto	C22 H21 N3 O2 S2	424
431) Line of the second se	C15 H19 N3 O2 S2	338
432) A Field	C16 H23 N3 O2 S2	354

Example	Structure	Molecular Formula	(M+H)+
433		C18 H19 N3 O2 S2	374
434		C18 H16 N4 O2 S2	385
435	Charles Charles	C20 H23 N3 O2 S2	402
436	A CI	C18 H17 F2 N3 O2 S2	410
437		C21 H23 N3 O2 S2	414
438		C18 H16 N4 O2 S3	417
439	and a	C19 H19 N3 O4 S2	- 418
440	3,01	C20 H23 N3 O3 S2	418
441	Lo - Chi	C18 H18 N4 O4 S2	419

Example :	Structure	Molecular Formula :	(M+H)+
442	it, sof	C18 H18 N4 O4 S2	419
443		C18 H18 N4 O4 S2	419
444		C19 H21 N3 O4 S2	420
445		C19 H21 N3 O4 S2	420
446	INH.	C18 H19 N5 O2 S3	434
447		C18 H19 N5 O2 S3	434
448	Z C C	C19 H18 F3 N3 O2 S2	442
449		C18 H18 Br N3 O2 S2	453
450	P. C.	C21 H25 N3 O5 S2	464

Example	Structure	Molecular Formula	(M+H)+
451	this to	C23 H28 N4 O4 S2	489
452	Of the second	C20 H21 N3 O2 S2	400
453	territy.	C18 H25 N3 O2 S2	380
454	5,424	C19 H21 N3 O2 S2	388
455	Start.	C27 H26 N4 O3 S2	519
456		C19 H21 N3 O3 S2	404
457	To the state of th	C18 H18 N4 O5 S2	435
458		C20 H23 N3 O2 S2	402

Example	Structure	Molecular Formula	(M+H)+
459		C24 H25 N3 O3 S2	468
460		C19 H21 N3 O2 S2	388
461	Creat Creat	C19 H21 N3 O2 S2	388
462	Com Com	C19 H21 N3 O3 S2	404
463		C17 H18 N4 O2 S2	489
464	De principal	C26 H28 N4 O4 S3	557
465) or	C19 H27 N3 O2 S2	394
466		C22 H22 N4 O3 S2	455

Example	Structure	Molecular Formula	(M+H)+
467		C22 H25 N3 O4 S2	460
468	5. 12. 12. 12. 12. 12. 12. 12. 12. 12. 12	C20 H21 N3 O3 S2	416
469) til tight	C15 H19 N3 O4 S2	370
470	Tri ta cori	C20 H18 F3 N3 O2 S2	454
471		C24 H26 N4 O3 S2	483
472	-Ora	C18 H19 N3 O3 S2	390
473		C18 H19 N3 O3 S2	390
474	La office of	C20 H20 N4 O2 S2	413

Example	Structure	Molecular Formula	- (M+H)+
475	Harry	C15 H21 N3 O2 S2	340
476	Christ Child	C19 H18 N4 O2 S2	. 399
477	Party.	C17 H18 N4 O2 S2	489
478	Start.	C17 H18 N4 O2 S2	489
479		C20 H20 N4 O2 S2	413
480	400	C20 H24 N4 O2 S2	531
481	raigi	C21 H22 N4 O2 S2	427
482	Jana de	C15 H17 N5 O2 S3	510

Example .	Structure	Molecular Formula	(M+H)+
483	3.72 C	C19 H21 N3 O3 S2	404
484	H. C.	C15 H17 N5 O2 S2	478
485	A.C.	C16 H17 N5 O4 S2	408
486	+3-10-1-10-1-10-1-10-1-10-1-10-1-10-1-10	C25 H32 N4 O4 S2	517
487	trore	C23 H28 N4 O4 S2	489
488	+10 T	C24 H30 N4 O4 S2	503
489		C19 H18 N6 O2 S3	459
490		C20 H24 N4 O2 S2	531

Example :	Structure :	Molecular Formula	(M+H)+
491	et of the state of	C18 H20 N4 O2 S2	503
492	C. Started	C19 H22 N4 O2 S2	517
493	Joseph North	C13 H18 N4 O2 S2	363
494	X () ()	C18 H18 F2 N4 O2 S2	425
495	X CHY	C18 H18 Cl2 N4 O2 S2	458
496		C17 H18 N4 O2 S2	489
497		C18 H20 N4 O2 S2	389
498	J. J	C14 H19 N3 O2 S2	326

Example	Structure	Molecular Formula	(M+H)+
499		C16 H21 N3 O2 S2	352
500		C14 H19 N3 O2 S2	326
501	X STORY	C14 H19 N3 O2 S2	326 -
502	OHON	C17 H17 N3 O3 S2	376
503	arrigh	C18 H19 N3 O3 S2	390
504	THE	C14 H19 N3 O3 S2	342
505	Chiral	C21 H31 N3 O3 S2	438
506	S S CONH,	C10 H9 Br N4 O3 S2	378
507		C19 H22 N4 O3 S2	419
508		C18 H20 N4 O2 S2	389

Example	Structure	Molecular Formula	(M+H)+
509		C19 H22 N4 O2 S2	403
510		C19 H22 N4 O2 S2	403
511	X S S S S S S S S S S S S S S S S S S S	C15 H21 N3 O3 S2	356
512	and the	C23 H27 N3 O2 S2	442
513	Xa	C21 H29 N3 O2 S2	420
514	Sta	C24 H25 N3 O2 S2	452
515	gr conto	C24 H25 N3 O2 S2	452
516	Xa	C17 H23 N3 O2 S2	. 366
517	Xª Fiel	C18 H27 N3 O2 S2	382

Example	Structure	Molecular Formula	(M+H)+
518		C20 H23 N3 O2 S2	402
519		C20 H20 N4 O2 S2	413
520	C. T. C. S.	C22 H27 N3 O2 S2	430
521	to the	C20 H21 F2 N3 O2 S2	438
522	S'HILLY	C23 H27 N3 O2 S2	442
523		C20 H20 N4 O2 S3	445
524	and of the second	C21 H23 N3 O4 S2	446
525	S. S	C22 H27 N3 O3 S2	446

Exc	mple	Structure	Molecular Formula	(M+H)+
	526	to still to	C20 H22 N4 O4 S2	447
	527	£ rof	C20 H22 N4 O4 S2	447
	528	X Y Y	C20 H22 N4 O4 S2	447
	529	C'hi	C21 H25 N3 O3 S2	·432
	530		C21 H25 N3 O4 S2	448
	531		C20 H23 N5 O2 S3	462
	532		C20 H23 N5 O2 S3	462
	533	Strip of	C21 H22 F3 N3 O2 S2	470
	534	X DiD	C20 H22 Br N3 O2 S2	481

Example i	Structure :	Molecular Formula	(M+H)+
535	The state of the s	C23 H29 N3 O5 S2	492
536		C21 H24 N4 O3 S2	445
537	E Harry	C22 H25 N3 O4 S2	. 460
538	Orgo Sax	C20 H29 N3 O2 S2	408
539	9. July 10. 4	C21 H25 N3 O2 S2	416
540	S. X	C29 H30 N4 O3 S2	547
541	S. Jarox	C22 H27 N3 O3 S2	446
542	10°	C20 H22 N4 O5 S2	463

Example	Structure	Molecular Formula	(M+H)+
543		C22 H27 N3 O2 S2	430
544		C26 H29 N3 O3 S2	496
545		C21 H25 N3 O2 S2	416
546	to by	C25 H32 N4 O4 S2	517
547	the first	C26 H34 N4 O4 S2	531
548	O Co	C19 H22 N4 O2 S2	517 ^
549	Xy City	C17 H21 N5 O4 S2	424
550	org org	C21 H31 N3 O2 S2	422

Example	Structure	Molecular Formula	(M+H)+
551		C24 H26 N4 O3 S2	483
552	چې د پې د	C24 H29 N3 O4 S2	488
553	3,p2	C22 H25 N3 O3 S2	444
554	, se	C21 H25 N3 O4 S2	448 [.]
555	S. S	C21 H25 N3 O3 S2	432
556		C26 H30 N4 O3 S2	511
557		C20 H23 N3 O3 S2	418
558	gy and	C20 H23 N3 O3 S2	418

Example	Structure :	Molecular Formula	(M+H)+
559	or or	C20 H23 N3 O3 S2	418
560		C20 H22 N4 O5 S2	463
561	Xª G.	C17 H25 N3 O2 S2	368
562	J. J	C20 H23 N3 O4 S2	434
563	or or	C19 H22 N4 O2 S2	517
564	O.C.	C19 H22 N4 O2 S2	517
565		C22 H24 N4 O2 S2	441
566	Joseph Sax	C22 H28 N4 O2 S2	559

· 5722

Example	Structure	Molecular Formula	(M+H)+
567	thair	C23 H26 N4 O2 S2	569
568	to the	C17 H21 N5 O2 S3	538
569	Start Start	C21 H25 N3 O3 S2	432
570	X	C17 H21 N5 O2 S2	506
571		C18 H21 N5 O4 S2	436
572	43. C.	C27 H36 N4 O4 S2	545
573	trora	C25 H32 N4 O4 S2	517
574	+40x 13	C26 H34 N4 O4 S2	531

Example	Structure	Molecular Formula	(M+H)+
575	CALLES TON	C21 H22 N6 O2 S3	487
576		C22 H28 N4 O2 S2	559
577	in the second se	C20 H24 N4 O2 S2	531
578	C. Jarox	C21 H26 N4 O2 S2	545
579	,OTO	C20 H24 N4 O2 S2	531
580	or ax	C21 H26 N4 O2 S2	545
581		C13 H15 N3 O4 S2	342
582	Joseph OH	C11 H13 N3 O3 S2	300

Example	Structure	Molecular Formula !	(M+H)+
583	N N N N N N N N N N N N N N N N N N N	C11 H14 N4 Ö2 S2 :	413
584		C17 H23 N3 O4 S2	398
585		C16 H21 N3 O4 S2	384
586	YYYY	C15 H21 N3 O3 S2	356
587		C18 H18 F2 N4 O3 S2	441
588		C18 H18 F2 N4 O4 S2	457
589	The state of	C15 H21 N3 O5 S2	388
590	You have a second	C15 H21 N3 O4 S2	372
591	J. J	C17 H17 N3 O3 S2	376
592	٥٠٠٠	C21 H22 Cl2 N4 O2 S2	498
593	d'hisimo	C21 H22 F2 N4 O2 S2	465

Example	Structure	Molecular Formula	(M+H)+
594		C14 H19 N3 O2 S2	326
595	OH S S OH	C10 H11 N3 O3 S2	286
596	D'I'S	C18 H19 F N4 O4 S2	439
597	Dirit	C18 H19 F N4 O2 S2	407
598 ·	philip	C18 H19 F N4 O3 S2	423
599		C15 H21 N3 O4 S2	372
600		C14 H19 N3 O3 S2	342
601		C14 H19 N3 O4 S2	358
602	+9,50	C14 H20 N4 O2 S2	341

Example i	Structure	Molecular Formula	(M+H)+
603	XII S-ÇIÎ	C18 H19 F N4 O2 S2	407
604	X - Ti	C18 H18 F2 N4 O2 S2	425
605	X-al	C18 H17 F3 N4 O2 S2	443
606		C18 H19 CI N4 O2 S2	423
607	Sarah.	C21 H26 N4 O2 S2	431
608	+9	C15 H22 N4 O3 S2	371
609	+57	C16 H24 N4 O3 S2	385
610	trat.	C19 H22 N4 O3 S2	419

Example :	Structure	Molecular Formula :	(M+H)+
611	tri ch	C19 H21 F N4 O3 S2	437
612	ara A	C19 H22 N4 O3 S2	419
613	w.	C19 H20 N4 O4 S2	433
614	4	C18 H27 N5 O2 S2	524
615	+ 1	C17 H22 N6 O2 S2	521
616	+9	C14 H17 N7 O2 S2	494
617	La La	C19 H21 N5 O3 S2	432
618	O'CO	C17 H19 N5 O2 S2	504

Example :	Structure	Molecular Formula	(M+H)+
619	Trib	C22 H25 N5 O2 S2	456
620	TAY D	C18 H24 N6 O2 S2	535
621		C21 H23 F N4 O2 S2	447
622		C21 H22 F2 N4 O2 S2	465
623		C21 H21 F3 N4 O2 S2	483
624		C21 H23 CI N4 O2 S2	464
625	Qa Gio	C24 H30 N4 O2 S2	471
626	Carrier.	C18 H26 N4 O3 S2	411

Example	Structure 1	Molecular Formula	(M+H)+
627	Q tratical	C19 H28 N4 O3 S2	425
628	Qui rio	C22 H26 N4 O3 S2	459
629	Sar di	C22 H25 F N4 O3 S2	477
630	Ga Car	C22 H26 N4 O3 S2	459
631	O'A Line	C22 H24 N4 O4 S2	473
632	Of the o	C21 H31 N5 O2 S2	564
633	Se str	C20 H26 N6 O2 S2	561
634	Ca tivo	C17 H21 N7 O2 S2	534

Example	Structure	Molecular Formula	(M+H)+
635	Qa firoi	C23 H29 N5 O2 S2	586
636	Qa Giror	C22 H25 N5 O3 S2	472
637	or de	C20 H23 N5 O2 S2	544
638	St. B	C25 H29 N5 O2 S2	496
639	Sept.	C21 H28 N6 O2 S2	575
640	140×4	C24 H33 N3 O3 S2 Si	504
641	-a-	C23 H28 N4 O4 S2	489
642	and a series	C19 H28 N4 O2 S2	409
643	HC CH S COBHS NO	C15 H21 N3 O2 S2	340

Example	Structure	Molecular Formula	(M+H)+
644	Mic Oil	C17 H23 N3 O2 S2	367
645	Jan Coron	C24 H31 N5 O2 S2	487
646	"The Ca	C19 H28 N4 O2 S2	410
647	Lord	C19 H28 N4 O2 S2	410
648	J. Olo	C18 H27 N5 O2 S2	411
649		C16 H19 N5 O2 S2	378
650	MC CH	C16 H18 N4 O S2	347
651	Michael State of the State of t	C17 H19 N3 O S2	346
652	"Z" TILL	C19 H22 N4 O2 S2	404
653		C19 H22 N4 O2 S2	404
654	" CILONO	C25 H32 N4 O3 S2	502
655	"X" TILL	C20 H24 N4 O2 S2	418

Example	Structure	Molecular Formula	(M+H)+
656	" C.	C19 H23 N4 O2 S2	405
657		C18 H20 N4 O3 S2	406
658	me Company	C16 H18 N4 O3 S2	379
659	MC PH	C16 H18 N4 O2 S2	363
660	S S COL	C16 H17 Br N4 O S2	426
661	ang	C19 H23 N3 O3 S2	407
662	30.00mg	C21 H30 N6 O S2	448
663	";> (1-1) (1)	C19 H25 N5 O2 S2	421
664		C17 H18 N4 O2 S2	375
665	in the second	C24 H31 N5 O3 S2	503
666	" ALON"	C21 H26 N4 O3 S2	448
667		C17 H20 N4 O2 S2	378

Example	Structure	Molecular Formula	(M+H)+
668	\$ Company	C21 H27 N5 O3 S2	463
669		C19 H23 N5 O3 S2	435
670	المراجعة الم	C15 H17 N5 O2 S2	364
671	"X" on the	C19 H22 N4 O2 S2	404
672	CN S S N N	C13 H11 N5 S2	302
673	Q ~ C ~ C	C14 H12 N4 S2	301
674	N,C SSYND	C17 H18 N4 S2	343
675	MC ST ST NO	C17 H18 N4 S2	343
676	H,C S S N N	C15 H14 N4 S2	3,15
677		C16 H18 N4 O2 S2	363
678		C16 H18 N4 O2 S2	363
679	">" O 10 10 10 10 10 10 10 10 10 10 10 10 10	C22 H31 N5 O2 S2	463

Example	Structure :	Molecular Formula	(M+H)+
680		C20 H24 N4 O4 S2	450
681	"\$ C.	C21 H27 N5 O S2	431
682	"HUOO	C21 H27 N5 O3 S2	463
683	STO CO	C22 H31 N5 O3 S2	479
684	St. OO	C21 H27 N5 O2 S2	447
685	graioù.	C23 H29 N3 O5 S2	493
686	3000	C23 H29 N3 O5 S2	493
687	" Sand	C22 H31 N5 O S2	447
688		C22 H28 N4 O2 S2	446
689	"% To	C20 H26 N4 O2 S2	420
690	" DLaron"	C22 H31 N5 O2 S2	463
691	"STATOO	C22 H28 N4 O3 S2	462

Example	Structure	Molecular Formula	(M+H)+
692	*franco	C25 H32 N4 O3 S2	502
693		C21 H25 N3 O4 S2	449
694	grows.	C20 H24 N4 O2 S2	418
695	Lesion	C25 H34 N4 O3 S2	504
696	a coraz	C24 H30 N4 O2 S2	472
697	a conste	C24 H30 N4 O3 S2	; ! . 488
698	-may st	C22 H28 N4 O3 S2	462
699	"Lange	C24 H33 N5 O2 S2	489
700	-rausse	C23 H28 N4 O4 S2	490
701	anautola	C26 H35 N5 O2 S2	515
702		C20 H23 N3 O3 S2	419
703	Lange	C43 H49 N7 O6 S4	889

Example	Structure 1	Molecular Formula	(M+H)+
704		C20 H23 N3 O4 S3	467
705	200000	C25 H32 N4 O4 S2	518
706		C17 H20 N4 O4 S3	442
707	"Sarano".	C21 H24 CI N3 O3 S2	467
708	*pranon	C22 H28 N4 O4 S2	478
709	2000°	C21 H26 N4 O3 S2	448
710	ga-arong.	C25 H32 N4 O5 S3	566
711	" and	C22 H28 N4 O5 S3	526
712	The State of the	C19 H22 N4 O4 S3	468
713	anox	C22 H28 N4 O3 S2	462
714	do de	C25 H34 N4 O3 S2	504
715	"LOW To	C22 H32 N4 O4 S2	482

Example	Structure	Molecular Formula	(M+H)+
716	The second	C17 H24 N4 O2 S2	382
717	The Contraction of the Contracti	C18 H26 N4 O4 S3	460
718	"CO CO CONTROL OF COM	C18 H26 N4 O2 S2	396
719	on of	C24 H33 N5 O2 S2	489
720	o corst	C26 H35 N5 O2 S2	515
721	S. O.L.	C24 H30 N4 O2 S2	. 472
722	E. O.F.	C20 H24 N4 O2 S2	418
723	op op	C24 H30 N4 O3 S2	488
724	arat .	C26 H38 N4 O2 S2	504
725	: Siiiiii	C23 H29 N5 O4 S2	505
726	"%" WILL	C25 H32 N4 O4 S2	518
727	Lymano	C25 H31 N5 O3 S2	515

Example	Structure	Molecular Formula	(M+H)+
728		C19 H25 N5 O3 S2	437
729	20-andre	C22 H32 N4 O4 S2	482
730	HC COL H,C S IN O	C17 H24 N4 O2 S2	382
731	The Contraction of the Contracti	C18 H26 N4 O2 S2	396
732	med of the state o	C18 H21 N5 O2 S2	405
733	Me St. Mess.	C18 H26 N4 O4 S3	460
734	"Y" OLLOWA	C24 H30 N4 O3 S2	488
735	-X-oro	C26 H36 N4 O4 S2	534

What is Claimed is:

1. A compound of the formula

$$R_3 \xrightarrow{R_1} S(O)_m \xrightarrow{S} \overset{H}{N} R_4$$
 (I)

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and pharmaceutically acceptable salts thereof wherein:

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

R₃ is aryl or heteroaryl;

R4 is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl,

10 heteroarylalkyl, heterocycloalkyl,

heterocycloalkylalkyl; or

CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl, CO-heteroaryl, CO-alkyl-heteroaryl, CO-heterocycloalkyl,

CO-alkyl-heterocycloalkyl; or

15 CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl,

CONH-alkyl-aryl, CONH-heteroaryl,

CONH-alkyl-heteroaryl, CONH-heterocycloalkyl,

CONH-alkyl-heterocycloalkyl; or

- COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl,
- COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl,

COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or

SO2-cycloalkyl, SO2-aryl, SO2-alkyl-cycloalkyl, SO2-alkyl-aryl,

SO₂-heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl,

SO₂-alkyl-heterocycloalkyl; or

25 C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl,

C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl,

C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl,

C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or

C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl,
C(NNO₂)NH-alkyl-cycloalkyl, C(NNO₂)NH-alkyl-aryl,
C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl,
C(NNO₂)NH-heterocyloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl;

5 or

C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl, C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl, C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl,

C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or

10 C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,
C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl,
C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl,
C(NH)NHCO-heterocylcloalkyl,

C(NH)NHCO-alkyl-heterocycloalkyl; or

- C(NOR₆)NH-alkyl, C(NOR₆)NH-cycloalkyl, C(NOR₆)NH-aryl,
 C(NOR₆)NH-alkyl-cycloalkyl, C(NOR₆)NH-alkyl-aryl,
 C(NOR₆)NH-heteroaryl, C(NOR₆)NH-alkyl-heteroaryl,
 C(NOR₆)NH-heterocylcoalkyl, C(NOR₆)NH-alkyl-heterocycloalkyl;
 R₅ is hydrogen or alkyl;
- 20 R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl; m is an integer of 0 to 2; and n is an integer of 1 to 3.
- The compounds as recited in Claim 1, wherein
 R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_3

wherein Y is oxygen, sulfur or NR9

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R4 is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl, heterocycloalkylalkyl; or CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl, CO-heteroaryl, CO-alkyl-heteroaryl, CO-heterocycloalkyl, 5 CO-alkyl-heterocycloalkyl; or CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl, CONH-alkyl-aryl, CONH-heteroaryl, CONH-alkyl-heteroaryl, CONH-heterocycloalkyl, 10 CONH-alkyl-heterocycloalkyl; or COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl, COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl, COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or SO2-cycloalkyl, SO2-aryl, SO2-alkyl-cycloalkyl, SO2-alkyl-aryl, SO₂-heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl, 15 SO₂-alkyl-heterocycloalkyl; or C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl, C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl, C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl, C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or 20 C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl, C(NNO2)NH-alkyl-cycloalkyl, C(NNO2)NH-alkyl-aryl, C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl, C(NNO₂)NH-heterocyloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl; 25 C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl, C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl, C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl, C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or

C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,

C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl,

C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl, C(NH)NHCO-heterocylcloalkyl,

C(NH)NHCO-alkyl-heterocycloalkyl; or

C(NOR6)NH-alkyl, C(NOR6)NH-cycloalkyl, C(NOR6)NH-aryl,

5 C(NOR6)NH-alkyl-cycloalkyl, C(NOR6)NH-alkyl-aryl,
C(NOR6)NH-heteroaryl, C(NOR6)NH-alkyl-heteroaryl,
C(NOR6)NH-heterocylcoalkyl, C(NOR6)NH-alkyl-heterocycloalkyl;

R₅ is hydrogen or alkyl;

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R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

R₇ and R₈ are independently hydrogen, alkyl, substituted alkyl, cycloalkyl, aryl, substituted aryl, cycloalkylalkyl, arylalkyl, heteroaryl, substituted heteroaryl, heteroarylalkyl, heterocycloalkyl, heterocycloalkylalkyl;

R₉ is hydrogen, alkyl, cycloalkyl, aryl, akylcycloalkyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl; m is an integer of 0 to 2; and

20 3. The compounds as recited in Claim 1, wherein

n is an integer of 1 to 3.

R₁ and R₂ are independently hydrogen, fluorine or alkyl;-

$$R_3$$
 is N_{N_2}

wherein Y is oxygen;

R4 is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl,

25 heteroarylalkyl, heterocycloalkyl,

heterocycloalkylalkyl; or

CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl, CO-heteroaryl, CO-alkyl-heteroaryl, CO-heterocycloalkyl, CO-alkyl-heterocycloalkyl; or

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CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl, CONH-alkyl-aryl, CONH-heteroaryl,

CONH-alkyl-heteroaryl, CONH-heterocycloalkyl,

CONH-alkyl-heterocycloalkyl; or

5 COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl, COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl,

COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or

SO₂-cycloalkyl, SO₂-aryl, SO₂-alkyl-cycloalkyl, SO₂-alkyl-aryl, SO₂-

heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl, SO₂-alkyl-

10 heterocycloalkyl; or

C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl,

C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl,

C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl,

C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or

15 C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl,

C(NNO2)NH-alkyl-cycloalkyl, C(NNO2)NH-alkyl-aryl,

C(NNO2)NH-heteroaryl, C(NNO2)NH-alkyl-heteroaryl,

C(NNO2)NH-heterocyloalkyl, C(NNO2)NH-alkyl-heterocycloalkyl;

or

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20 C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl,

C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl,

C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl,

C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or

C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,

C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl,

C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl,

C(NH)NHCO-heterocylcloalkyl,

C(NH)NHCO-alkyl-heterocycloalkyl; or

C(NOR₆)NH-alkyl, C(NOR₆)NH-cycloalkyl, C(NOR₆)NH-aryl,

30 C(NOR₆)NH-alkyl-cycloalkyl, C(NOR₆)NH-alkyl-aryl,

C(NOR₆)NH-heteroaryl, C(NOR₆)NH-alkyl-heteroaryl,

3,----

C(NOR₆)NH-heterocylcoalkyl, C(NOR₆)NH-alkyl-heterocycloalkyl; R₅ is hydrogen;

R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

R₇ and R₈ are independently hydrogen, alkyl, substituted alkyl, cycloalkyl, aryl, substituted aryl, cycloalkylalkyl, arylalkyl, heteroaryl, substituted heteroaryl, heteroarylalkyl, heterocycloalkyl, heterocycloalkylalkyl;

m is an integer of 0 to 2; and n is an integer of 1 to 3.

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4. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is $N_{N_2} = R_8$

wherein Y is sulfur;

R₄ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl,

heterocycloalkylalkyl; or

CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl,

 $CO\text{-}heteroaryl, CO\text{-}alkyl\text{-}heteroaryl, CO\text{-}heterocycloalkyl,}$

CO-alkyl-heterocycloalkyl; or

CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl,

CONH-alkyl-aryl, CONH-heteroaryl,

CONH-alkyl-heteroaryl, CONH-heterocycloalkyl,

CONH-alkyl-heterocycloalkyl; or

COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl,

COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl,

COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or

SO2-cycloalkyl, SO2-aryl, SO2-alkyl-cycloalkyl, SO2-alkyl-aryl,

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SO_2-heteroaryl, SO_2-alkyl-heteroaryl, SO_2-heterocycloalkyl, SO_2-alkyl-heterocycloalkyl; or
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C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl,

C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl,

C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl,

C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or

C(NNO2)NH-alkyl, C(NNO2)NH-cycloalkyl, C(NNO2)NH-aryl,

C(NNO₂)NH-alkyl-cycloalkyl, C(NNO₂)NH-alkyl-aryl,

C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl,

C(NNO₂)NH-heterocyloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl;

or

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C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl,

C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl,

C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl,

C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or

C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,

C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl,

C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl,

C(NH)NHCO-heterocylcloalkyl,

20 C(NH)NHCO-alkyl-heterocycloalkyl; or

C(NOR₆)NH-alkyl, C(NOR₆)NH-cycloalkyl, C(NOR₆)NH-aryl,

C(NOR6)NH-alkyl-cycloalkyl, C(NOR6)NH-alkyl-aryl,

C(NOR6)NH-heteroaryl, C(NOR6)NH-alkyl-heteroaryl,

C(NOR6)NH-heterocylcoalkyl, C(NOR6)NH-alkyl-heterocycloalkyl;

25 R₅ is hydrogen;

R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

 R_7 and R_8 are independently hydrogen, alkyl, substituted alkyl,

cycloalkyl, aryl, subsituted aryl, cycloalkylalkyl, arylalkyl, heteroaryl,

substituted heteroaryl, heteroarylalkyl, heterocycloalkyl, heterocycloalkylalkyl;

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m is an integer of 0 to 2; and n is an integer of 1 to 3.

5. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_7

wherein Y is NR₉;

R₄ is alkyl, cycloalkyl, aryl, cycloalkylalkyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl,

10 heterocycloalkylalkyl; or

CO-alkyl, CO-cycloalkyl, CO-aryl, CO-alkyl-cycloalkyl, CO-alkyl-aryl, CO-heteroaryl, CO-heterocycloalkyl, CO-alkyl-heterocycloalkyl; or

CONH-alkyl, CONH-cycloalkyl, CONH-aryl, CONH-alkyl-cycloalkyl, CONH-alkyl-aryl, CONH-heteroaryl, CONH-heterocycloalkyl, CONH-alkyl-heterocycloalkyl; or

COO-alkyl, COO-cycloalkyl, COO-aryl, COO-alkyl-cycloalkyl, COO-alkyl-aryl, COO-heteroaryl, COO-alkyl-heteroaryl,

COO-heterocycloalkyl, COO-alkyl-heterocycloalkyl; or SO₂-cycloalkyl, SO₂-aryl, SO₂-alkyl-cycloalkyl, SO₂-alkyl-aryl,

SO₂-heteroaryl, SO₂-alkyl-heteroaryl, SO₂-heterocycloalkyl, SO₂-alkyl-heterocycloalkyl; or

C(NCN)NH-alkyl, C(NCN)NH-cycloalkyl, C(NCN)NH-aryl,

25 C(NCNNH)-alkyl-cycloalkyl, C(NCN)NH-alkyl-aryl, C(NCN)NH-heteroaryl, C(NCN)NH-alkyl-heteroaryl,

C(NCN)NH-heterocycloalkyl, C(NCN)NH-alkyl-heterocylcoalkyl; or

C(NNO₂)NH-alkyl, C(NNO₂)NH-cycloalkyl, C(NNO₂)NH-aryl, C(NNO₂)NH-alkyl-cycloalkyl, C(NNO₂)NH-alkyl-aryl,

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C(NNO₂)NH-heteroaryl, C(NNO₂)NH-alkyl-heteroaryl, C(NNO₂)NH-heterocycloalkyl, C(NNO₂)NH-alkyl-heterocycloalkyl;

C(NH)NH-alkyl, C(NH)NH-cycloalkyl, C(NH)NH-aryl,

5 C(NH)NH-alkyl-cycloalkyl, C(NH)NH-alkyl-aryl,

C(NH)NH-heteroaryl, C(NH)NH-alkyl-heteroaryl,

C(NH)NH-heterocycloalkyl, C(NH)NH-alkyl-heterocycloalkyl; or

C(NH)NHCO-alkyl, C(NH)NHCO-cycloalkyl, C(NH)NHCO-aryl,

C(NH)NHCO-alkyl-cycloalkyl, C(NH)NHCO-alkyl-aryl,

C(NH)NHCO-heteroaryl, C(NH)NHCO-alkyl-heteroaryl,

C(NH)NHCO-heterocylcloalkyl,

C(NH)NHCO-alkyl-heterocycloalkyl; or

C(NOR₆)NH-alkyl, C(NOR₆)NH-cycloalkyl, C(NOR₆)NH-aryl,

C(NOR6)NH-alkyl-cycloalkyl, C(NOR6)NH-alkyl-aryl,

C(NOR₆)NH-heteroaryl, C(NOR₆)NH-alkyl-heteroaryl,

C(NOR₆)NH-heterocylcoalkyl, C(NOR₆)NH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

or

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R₆ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl;

R₇ and R₈ are independently hydrogen, alkyl, substituted alkyl, cycloalkyl, aryl, substituted aryl, cycloalkylalkyl, arylalkyl, heteroaryl, substituted heteroaryl, heteroarylalkyl, heterocycloalkyl, heterocycloalkyl;

R₉ is hydrogen, alkyl, cycloalkyl, aryl, cycloalkylakyl, arylalkyl, heteroaryl, heteroarylalkyl, heterocycloalkyl or heterocycloalkylalkyl; m is an integer of 0 to 2; and

n is an integer of 1 to 3.

6. The compounds as recited in Claim 1, wherein

30 R₁ and R₂ are independently hydrogen, fluorine or alkyl;

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$$R_3$$
 is N_{N_2}

wherein Y is oxygen;

R4 is CO-alkyl, CO-alkyl-aryl, CO-cycloalkyl, CO-alkyl-heteroaryl, CO-alkyl-heteroalkyl, CO-alkyl-heterocycloalkyl, aryl, arylalkyl,

5 heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen; and

R7 and R8 are hydrogen;

m is the integer 0; and

n is the integer 1.

7. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

wherein Y is oxygen;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

20 R₅ is hydrogen;

 R_7 and R_8 are alkyl;

m is the integer 0; and

n is the integer 1.

25 8. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is N

wherein Y is oxygen;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl,

5 heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is hydrogen;

R₈ is alkyl;

m is the integer 0; and

n is the integer 1.

9. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_7

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wherein Y is oxygen;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, CONH-alkyl,

20 CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is alkyl;

R₈ is hydrogen;

m is the integer 0; and

25 n is the integer 1.

10. The compounds as recited in Claim 1, wherein

 R_1 and R_2 are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_7

wherein Y is sulfur;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heterocycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl,

5 heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is hydrogen;

R₈ is alkyl;

m is the integer 0; and

n is the integer 1

11. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is $N_{N_2} = R_1$

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wherein Y is sulfur;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heterocycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, CONH-alkyl,

20 CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is alkyl;

R₈ is hydrogen;

m is the integer 0; and

25 n is the integer 1.

12. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is N_{N_2}

wherein Y is NR9;

R4 is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl,

5 heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is hydrogen;

R₈ is alkyl;

R₉ is hydrogen, alkyl, cycloalkyl, aryl, alkyl-cycloalkyl, alkyl-aryl, heteroaryl, alkyl-heteroaryl, heterocycloalkyl, or alkyl-heterocycloalkyl; m is the integer 0; and

n is the integer 1.

15 13. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_8

wherein Y is NR₉;

R4 is CO-alkyl, CO-alkyl-aryl, CO-alkyl-heteroalkyl, CO-cycloalkyl,

20 CO-alkyl-heterocycloalkyl, CO-alkyl-heteroaryl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

R₅ is hydrogen;

R₇ is alkyl;

25 R₈ is hydrogen;

R₉ is alkyl;

m is the integer 0; and

n is the integer 1.

14. The compounds as recited in Claim 1, wherein

R₁ and R₂ are independently hydrogen, fluorine or alkyl;

$$R_3$$
 is R_7

5 wherein X is NR₉;

R₄ is CO-alkyl, CO-alkyl-aryl, CO-cycloalkyl, CO-alkyl-heteroaryl, CO-alkyl-heteroalkyl, CO-alkyl-heterocycloalkyl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, CONH-alkyl,

CONH-alkyl-aryl, CONH-cycloalkyl, or CONH-alkyl-heterocycloalkyl;

10 R₅ is hydrogen;

R₇ is alkyl;

R₈ is hydrogen;

R₉ is hydrogen;

m is the integer 0

n is the integer 1.

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15. The compound as recited in Claim 1, which is

N-[5-[[5-Ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide;

N-[5-[[5-Ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]benzamide;

N-[5-[[(4,5-Dimethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide;

N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide;

N-[5-[[5-t-Butyl-2-oxazolyl)methyl]thio]-2-

thiazolyl]trimethylacetamide:

N-[5-[[(4-Ethyl-2-oxazolyl)methyl]thio]-2-thiazolyl]acetamide;

N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-N'-cyano-N"-(2,6-difluorophenyl)guanidine;

N-[5-[(5-Isopropyl-2-oxazolyl)fluoromethyl]thio]-2-thiazolyl acetamide;

N-[5-[[(5-t-Butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]

30 aminophenyl-4-(2-hydroxyethyl)sulfonamide;

N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]aminophenyl-4-sulfonamide;

N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-4-aminopyrimidine;

N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-3-(hydroxymethyl)aniline;

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N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2-thiazolyl]-2-aminopyridine;

N-[5-[[(5-t-butyl-2-oxazolyl)methyl]thio]-2- thiazolyl]-2-[5-[(((3-10 hydroxy-2,2-dimethyl)propyl)amino)methyl]] aminopyridine; or a pharmaceutically acceptable salt thereof.

- 16. A pharmaceutical composition comprising a compound of Claim 1 and a pharmaceutically acceptable carrier.
- 17. A pharmaceutical composition comprising a compound of Claim 1, in combination with a pharmaceutically acceptable carrier, and an anti-cancer agent formulated as a fixed dose.
- 20 18. A pharmaceutical composition comprising a compound of Claim 1, in combination with a pharmaceutically acceptable carrier, and a modulator of p53 transactivation formulated as a fixed dose.
- 19. A pharmaceutical composition according to claim 16, comprising a
 25 compound of Claim 1 in combination with a pharmaceutically acceptable carrier, with an anticancer treatment or anticancer agent administered in sequence.
- 20. The pharmaceutical composition according to Claim 18, wherein said combination comprising said compound of Claim 1 and said

pharmaceutically acceptable carrier, is administered prior to administration of said anticancer treatment or anticancer agent.

- 21. The pharmaceutical composition according to claim 18, wherein said combination comprising said compound of Claim 1 and said pharmaceutically acceptable carrier, is administered after administration of said anticancer treatment or anticancer agent.
- 22. A method of inhibiting protein kinases which comprises administering
 to a mammalian specie in need thereof an effective protein kinase inhibiting amount of a compound of Claim 1.
 - 23. A method of inhibiting cyclin dependent kinases which comprises administering to a mammalian specie in need thereof an effective cyclin dependent kinase inhibiting amount of a compound of Claim 1.
 - 24. A method of inhibiting cdc2 (cdk1) which comprises administering to a mammalian specie in need thereof an effective cdc2 inhibiting amount of a compound of Claim 1.

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- 25. A method of inhibiting cdk2 which comprises administering to a mammalian specie in need thereof an effective cdk2 inhibiting amount of a compound of Claim 1.
- 26. A method of inhibiting cdk3 which comprises administering to a mammalian specie in need thereof an effective cdk3 inhibiting amount of a compound of Claim 1.
- 27. A method of inhibiting cdk4 which comprises administering to a30 mammalian specie in need thereof an effective cdk4 inhibiting amount of a compound of Claim 1.

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28. A method of inhibiting cdk5 which comprises administering to a mammalian specie in need thereof an effective cdk5 inhibiting amount of a compound of Claim 1.

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- 29. A method of inhibiting cdk6 which comprises administering to a mammalian specie in need thereof an effective cdk6 inhibiting amount of a compound of Claim 1.
- 30. A method of inhibiting cdk7 which comprises administering to a mammalian specie in need thereof an effective cdk7 inhibiting amount of a compound of Claim 1.
- 31. A method of inhibiting cdk8 which comprises administering to a
 15 mammalian specie in need thereof an effective cdk8 inhibiting amount of a compound of Claim 1.
 - 32. A method for treating proliferative diseases comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.
 - 33. A method for treating cancer comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.

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34. A method for treating inflammation, inflamatory bowel disease, or transplantation rejection, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.

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35. A method for treating arthritis comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.

- 36. A method for treating infection by HIV, or for treating and preventing the development of AIDS, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.
- 10 37. A method for treating viral infections, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.
- 38. A method for treating fungal infections, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.
- 39. A method for preventing the development of cancer or tumor relapse,
 comprising administering to a mammalian specie in need thereof a
 therapeutically effective amount of a composition of Claim 16.
 - 40. A method for treating neurodegenerative disease, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 16.

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41. A method for treating proliferative diseases comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 17.

42. A method for treating cancer comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 17.

- 5 43. A method for preventing the development of cancer or tumor relapse, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 17.
- 44. A method for treating proliferative diseases comprising administering
 10 to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 18.
 - 45. A method for treating cancer comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 18.

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46. A method for preventing the development of cancer or tumor relapse, comprising administering to a mammalian specie in need thereof a therapeutically effective amount of a composition of Claim 18.

INTERNATIONAL SEARCH REPORT

Int anal Application No PCT/US 00/33037

A. CLASSI IPC 7	FICATION OF SUBJECT MATTER C07D277/54 C07D417/12 C07D417/	/14 A61K31/427 A61	P35/00					
According to	o International Patent Classification (IPC) or to both national classific	ation and IPC						
B. FIELDS SEARCHED								
Minimum do IPC 7	cumentation searched (classification system followed by classification CO7D A61K A61P	on symbols)						
	ion searched other than minimum documentation to the extent that s							
	ata base consulted during the international search (name of data ba BS Data, BEILSTEIN Data, WPI Data, E		ed)					
C. DOCUMI	ENTS CONSIDERED TO BE RELEVANT		****					
Category *	Citation of document, with indication, where appropriate, of the rel	evant passages	Relevant to claim No.					
X	WO 99 24416 A (BRISTOL-MYERS SQUI COMPANY) 20 May 1999 (1999-05-20) the whole document		1-46					
Furti	ner documents are listed in the continuation of box C.	X Patent family members are list	ed in annex.					
* Special ca *A* docume consid *E* earlier of filing d *L* docume which citation *O* docume other r *P* docume later th	International filing date with the application but theory underlying the se claimed invention not be considered to document is taken alone e claimed invention inventive step when the more other such document to a person stilled and tarnity search report							
9	March 2001	. ಕಿನಬಹಾ						
Name and n	nailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Authorized officer Alland, M						

INTERNATIONAL SEARCH REPORT

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